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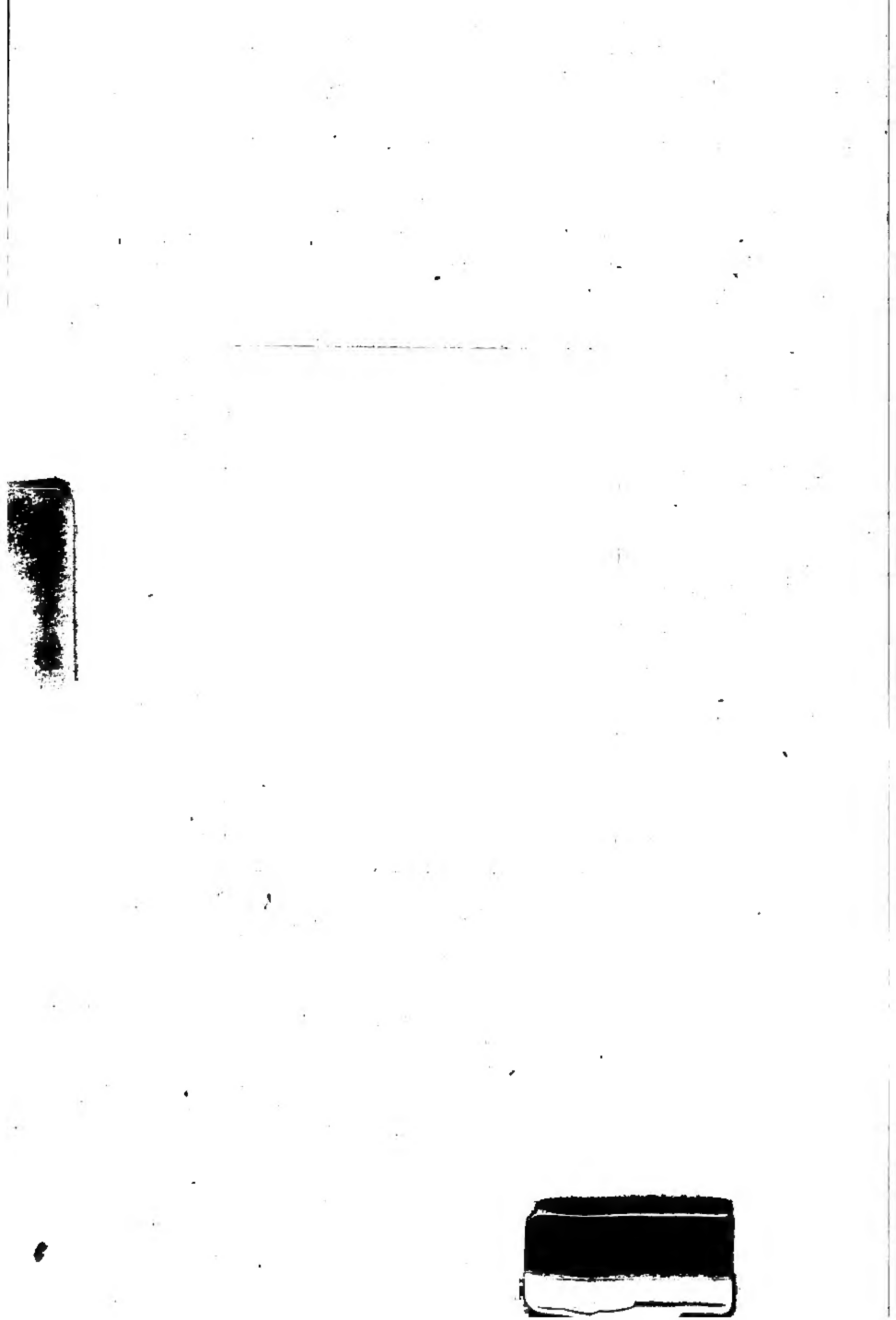
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FOOD INSPECTION AND ANALYSIS.

FOR THE USE OF PUBLIC ANALYSTS, HEALTH
OFFICERS, SANITARY CHEMISTS,
AND FOOD ECONOMISTS.

BY

ALBERT E. LEACH, S.B.,

Chief of the Denver Food and Drug Inspection Laboratory, Bureau of Chemistry, U. S. Department of Agriculture; formerly Chief Analyst of the Massachusetts State Board of Health.



SECOND EDITION, REVISED AND ENLARGED.

FIRST THOUSAND.

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Affectionately Dedicated to the Memory of Charles Pomeroy Worcester,

FORMER ANALYST OF THE MASSACHUSETTS STATE BOARD OF HEALTH,

WHOSE LOVABLE PERSONALITY AND STERLING INTEGRITY WERE

A CONSTANT INSPIRATION DURING MANY YEARS

OF CLOSE COMPANIONSHIP TO

THE AUTHOR.

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PREFACE TO SECOND EDITION.

During the five years that have elapsed since the appearance of the first edition, much progress has been made in food control work both in America and in Europe. In the United States the passage of the national pure food law, perhaps more than any other single factor, has contributed toward this, and has itself been the direct cause of ncreased activity on the part of many of the States. New standards have been adopted, many new methods have been tried out and found useful, and in some cases old ones have been displaced.

The most important of these changes and improvements have, it is believed, been embodied in the present edition, and include new material and modern methods of analysis covering a wide variety of subjects. Notable among these are meats and meat extracts, flour (including methods for determining the grade and for the detection of bleaching) noodles and Italian pastes, paprika, prepared mustard, tea, coffee, cocoa products (including milk chocolate), ice cream, maple products, honey, oils (including the Polenske number and Bömer's phytosterol-acetate test for vegetable oils), distilled liquors, preservatives (notably benzoic acid), etc.

A separate chapter on the refractometer, its varieties and application to food analysis has been introduced; also a separate chapter on flavoring extracts, including the lesser used extracts of almond, peppermint, wintergreen, rose, cassia, and cloves.

At the time the importance of a new edition seemed specially manifest, the author's health was such that it would have been impossible for him to personally undertake the work, and had it not been for his friends it could not have been accomplished. Indeed, the work of revision has been due to the untiring energy of Dr. A. L. Winton, Chief of the U. S. Food and Drug Inspection Laboratory at Chicago, who out of a busy life has taken entire charge of the details of the task.

supplying most of the new material, as well as introducing much that is original as a result of his ripe experience. To him, therefore, above all others, the author here expresses his deep appreciation and gratitude.

Special thanks are also extended to Dr. W. D. Bigelow, Chief of the Division of Foods of the Bureau of Chemistry, Washington, for his substantial work in revising the chapter on flesh foods, which includes much of his recent research along this line; also to Dr. T. B. Osborne, Chairman of the Committee on Protein Nomenclature of the American Physiological Society and of the Society of Biological Chemists, who has revised the classification of nitrogenous bodies; and finally to Mr. E. J. Shanley for his help in reading proof.

DENVER, COLORADO, September, 1909.

PREFACE TO FIRST EDITION.

In the preparation of the present work, the requirements of the public analyst are mainly kept in view, as well as of such officials as naturally cooperate with him in carrying out the provisions of the laws dealing with the suppression of food adulteration in states and municipalities. To this end special prominence is given to the nature and extent of adulteration in the various foods, to methods of analysis for the detection of adulterants, and to some extent also to the machinery of inspection.

While the analyst may not in all cases have directly to deal with the *minutiæ* of food inspection, his work is so closely allied therewith that this branch of the subject is of vital interest and importance to him. Indeed, in many smaller cities one official often has charge of the entire work, combining the duties of both inspector and analyst.

Endeavor has been made, furthermore, to deal with the general composition of foods, and to give such analytical processes as are likely to be needed by the sanitary chemist, or by the student who wishes to determine the proximate components of food materials.

It has been thought best to include brief synopses of processes of manufacture or preparation of certain foods and food materials, in cases where impurities might be suggested incidental to their preparation.

In view of the fact that Massachusetts was the pioneer state to adopt, over twenty years ago, a practical system of food and drug inspection, and for many years was the only state to enjoy such a system, no apology is perhaps needed for more frequent mention of Massachusetts methods and customs than those of many other states, in which the food laws are now being enforced with equal zeal and efficiency.

Considerable attention has been paid in the following pages to the use of the microscope in food analysis. Of the figures in the text illus-

viii PREFACE.

trating the microscopical structure of powdered tea, coffee, cocoa, and the spices, fifteen have been reproduced from the admirable drawings of Dr. Josef Moeller, of the University of Graz, Austria. Acknowledgment is gratefully given Dr. Moeller for his kind consent to their use.

The photomicrographs in half-tone, forming the set of plates at the end of the volume, were all made in the author's laboratory, and may be divided into three classes: 1st, illustrations of powdered pure foods and food products, as well as of powdered adulterants; 2d, types of adulterated foods, chosen from samples collected from time to time in the routine course of inspection; and 3d, photographs of permanently mounted sections of foods and adulterants.

While recent works covering the whole field of general food analysis are comparatively few, the number of treatises, monographs, government bulletins, and articles scattered through the journals, dealing with special subjects relative to food and its inspection, is surprisingly large, and from a painstaking review of these much information has been culled, for which it has been the author's intention at all times to give credit.

Special mention should here be made of the valuable publications of the U. S. Department of Agriculture, both the bulletins issued from Washington, and those from the various experiment stations, an everincreasing number of which are becoming engaged in human food work. The author has freely drawn from these sources, and especially from the data and material furnished by his coworkers in the recent and still pending labor of preparing food methods for the Association of Official Agricultural Chemists, and he wishes to extend his thanks to all of them for their assistance. Appreciation is also expressed for the care and discrimination shown by Mr. L. L. Poates in the preparation of the cuts. Thanks are especially due to Mr. Hermann C. Lythgoe, Assistant Analyst of the Massachusetts State Board of Health, for his invaluable cooperation, and to Dr. Thomas M. Drown for helpful hints and suggestions.

Boston, Mass., July 1, 1904.

TABLE OF CONTENTS.

CHAPTER I.
FOOD ANALYSIS AND OFFICIAL CONTROL
CHAPTER II.
THE LABORATORY AND ITS EQUIPMENT
CHAPTER III.
Food, its Functions, Proximate Components, and Nutritive Value 39-52 Nature and General Composition of Food; Fats, 39. Protein, and Classification of Nitrogenous Bodies, 40. Proteins, their Subdivisions, Occurrence, and Characterstic Tests, 40-45. Amino Acids, etc., 45. Alkaloids; Nitrates; Ammonia; Lecithin; Carbohdyrates and their Classification, 46. Organic Acids; Mineral or Inorganic Materials; Fuel Value of Food; Bomb Calorimeter, 47-48. References on Dietetics and Economy of Food, 49.
CHAPTER IV.
GENERAL ANALYTICAL METHODS

Determination of Free Ammonia; Determination of Amido Nitrogen, 74. Determination of Carbohydrates, 74. Poisoned Foods, 74. Detection and Determination of Arsenic, 75–76. Colorometric Analysis, 77. Tintometer, 78.

References on General Food Analysis, 79.

CHAPTER V.

THE MICROSCOPE IN FOOD ANALYSIS.

81-99
Microscopical vs. Chemical Analysis, 81. Technique of Food Microscopy, 82. Apparatus and Accessories, 82-84. Preparation of Vegetable Foods for Microscopical Examination, 85. Miscroscopical Diagnosis, 86. Vegetable Tissues and Cell Contents, under the Microscope, 87-90. Microscopical Reagents, 90-93. Microchemical Reactions, 90-93. Photomicrography; Appurtenances and Methods, 93-98.

References on the Microscope in Food Analysis, 98.

CHAPTER VI.

THE REFRACTOMETER.

Butyro-refractometer, 101. Refractometer Heater, 102. Manipulation, 102-104. Equivalents of Refractive Indices and Butyro-refractometer Readings, 105-106. Temperature Correction, 107. Abbé Refractometer, 108. Construction; Manipulation, 109-111. Immersion Refractometer, 111-112. Manipulation, 113-115. Equivalents of Refractive Indices and Immersion Refractometer Readings, 116-119. Strength of Solutions by Refractometer 120. Temperature Corrections, 121.

References on the Refractometer, 122.

CHAPTER VII.

Milk Adulteration and Inspection; Milk Standards, 159–161. Forms of Adulteration, and Variation in Standard, 161–162. Rapid Approximate Methods of Examination, 163–164. Examination of Milk Serum; Constants, 164–168. Systematic Routine Examination, 168. Analytical Methods for Solids, Fat, and Ash, 170–173. Added Foreign Ingredients, 173. Coloring Matters and their Detection, 174–177. Preservatives, their Relative Efficiency

and their Detection, 177-185. Added Cane Sugar, and Starch, 185. Added Condensed Milk; Analysis of Sour Milk, 186.

Condensed Milk; Composition, Standards, Adulteration, 186-188. Methods of Analysis, 188-191. Calculation of Fat in Original Milk, 192.

Cream; Composition, Analytical Methods, Standards, Adulterants, 193-195. Gelatin in Cream, 195-196. Sucrate of Lime in Cream, 196-198.

Ice Cream; Standard, Fillers, 198–199. Analytical Methods, 199–201. Cheese; Composition, Varieties, 201–202. Standards; Adulteration, 203–204. Analytical Methods, 188–191. Separation and Determination of Nitrogenous Bodies, 205–206. Lactic Acid; Milk Sugar; Foreign Fat, 207. References on Milk and its Products, 208.

CHAPTER VIII.

Meat; Structure and Composition, 211. Proximate Components of the Common Meats, 212-217. Meat Inspection, 217. Standards, 218. Meat Preservatives, 218. Curing, 219. Use of Antiseptics; Effect of Cooking, 220. Canned Meats, 221. Sausages, 223-224. Analytical Methods, 225. Fats of Meats, 226-227. Classification, Separation, and Determination of Nitrogenous Bodies, 228-231. Determination of Gelatin, 231. Determination of Nitrates, 232. Preservatives and their Detection, 232. Starch in Sausages, 233. Horseflesh in Sausages, and its Detection, 234-238. Muscle Sugar, 238. Coloring Matters and their Detection, 238-239. Detection of Frozen Meat, 239.

Meat Extracts; Character and Standards, 240–241. Composition, 242–244. Meat Juices, 245. Miscellaneous Meat Preparations, 246. Methods of Analysis, 246–249. Separation of Nitrogenous Compounds, 249–253. Acidity, 253. Preservatives; Glycerol, 254.

Fish; Structure Composition, and Methods of Analysis, 254-255. Crustaceans and Mollusks, 256. Analytical Methods; Preservatives in Fish and Oysters, 257.

Concentrated Foods for Armies and Campers, 257. References on Flesh Foods, 258.

CHAPTER IX.

Nature and Composition, 261. The Egg White and its Nitrogenous Compounds, 262. Preparation of Albumin; The Egg Yolk and its Composition, 263. Composition of the Ash, 264. Analytical Methods; Determination of Lecithin, 265. Preservation of Eggs, 266. Cold Storage Eggs, 267. Physical Methods of Examination, 267. Opened Eggs; Desiccated Eggs, 268. Egg Substitutes, 269. Custard Powders, 270.

References on Eggs, 270.

CHAPTER X.

PAGE

CEREALS AND THEIR PRODUCTS, LEGUMES, VEGETABLES, AND FRUITS...... 271-364
Composition of Cereals, Vegetables, Fruits, and Nuts, 271-276. Methods
of Proximate Analysis, 276-279. Carbohydrates of Cereals, 279. Starch;
Detection, Varieties, Classification, Microscopical Examination, 279-283.
Starch Determination, by Direct Acid Conversion and by Diastase Methods,
283-284. Cellulose; Crude Fiber, 285. Pentosans and their Determination,
285-294. Separation and Determination of the Carbohydrates of Cereals, 295296. Proteins of Cereals and Vegetables; Separation and Methods of Analysis,
296-298. Proteins of Wheat, their Separation and Determination, 298-300.
Proteins of Other Cereals and Vegetables, 300-301. Ash of Cereals and
Vegetables; Scheme for Ash Analysis, 301-305. Microscopy of Cereal Products, 305-311.

Flour; Milling, 311. Composition, 312. Damaged Flour; Ergot, 313. Adulteration, 314. Alum; Bleaching, 315. Inspection and Analysis; Fineness, 316. Pekar's Color Test; Absorption and Dough Test; Expansion of Dough, 317. Baking Tests, 317–319. Proximate Constituents; Gluten, 319. Protein; Acidity, 320. Detection of Bleaching; Nitrites, 321. Bamihl Gluten

Test, 322.

Bread; Composition; Varieties, 323-325. Methods of Examination, 325-326. Adulteration of Bread; Alum, 326. Cake, 327.

Leavening Materials; Yeast, 327. Compressed Yeast; Dry Yeast, 328. Composition and Microscopical Examination, 329. Yeast Testing; Available

Carbon Dioxide, 330. Starch in Compressed Yeast, 331.

Chemical Leavening Materials; Baking Powders, their Classification and Composition, 332-334. Adulteration, 334. Cream of Tartar and its Adulteration, 335. Analysis of Baking Chemicals, 336. Carbon Dioxide, 336-339. Tartaric Acid, 339-343. Starch, 343. Aluminum Salts, 344. Other Ingredients, 345-346.

Semolina, Macaroni, and Edible Pastes; Noodles, 347-348. Adulteration; Analytical Methods; Lecithin-Phosphoric Acid, 349. Colors, 349-

352. Shredded Wheat, 352.

Prepared Cereal Breakfast Foods; Nature and Composition, 352-354.

Analytical Methods, 354.

Infants' and Invalids' Foods, 354. Classification, 355. Composition, 356. Diabetic Foods, 357-358. Analytical Methods, 359-360.

References on Cereals, Vegetables, etc., 361. References on Leavening Materials, 364.

CHAPTER XI.

Coffee; Nature, Composition, Effect of Roasting, 379-381. Substitutes and Adulterants, 382. Analytical Methods; Caffetanic Acid, 382-383.

Caffeine, 384. Adulteration; Imitation Coffee; Coloring, 384. Glazing; Methods, 385. Microscopical Examination, 386. Chicory; its Microscopical Structure, 386-388. Composition of Chicory, and its Determination in Coffee,

389. Date Stones; Hygienic Coffee; Substitutes, 390-392.

Cocoa and Cocoa Products; Composition, Methods of Manufacture, 392-395. Theobromine and Nitrogenous Substances, 396. Milk Chocolate; Compounds, 397. Analytical Methods, 398. Starch; Sucrose; Lactose, 399. Theobromine and Caffeine, 400-401. Adulteration, and Standards of Purity, 402. Addition of Alkali, Microscopical Structure, 403-404. Cocoa Shells; Added Starch, Sugar, Fat and Colors, 405.

References on Tea, Coffee, and Cocoa, 406.

CHAPTER XII.

Methods of Proximate Analysis Common to all the Spices, 408. Moisture: Ash; Ether, and Alcohol Extract; Nitrogen; Starch; Crude Fiber; Volatile Oils, 400-411. Microscopical Examination, 412. Spice Adulterants, 412-413. Cloves; Composition, 412-415. Tannin, 415. Microscopical Examination.

416. Clove Stems, 417. Adulteration and Standard of Purity; Exhausted Cloves, 418. Cocoanut Shells, 419.

Allspice; Composition, 420. Tannin Equivalent, 421. Microscopical Structure, 422-423. Adulteration and Standard of Purity, 424.

Cassia and Cinnamon; Composition, 424-425. Microscopical Structure, 426-427. Adulterants; Standard, 428. Foreign Bark, 428.

Pepper; Composition, 428-432. Nitrogen Determination, 432. Piperin, 433. Microscopical Examination, 433-434. Adulteration and Standards, 435. Pepper Shells and Dust, 435. Olive Stones, 436. Buckwheat, 437. Long Pepper, 438.

Red Pepper; (Cayenne, Paprika, etc.). Nature; Varieties: Composition, 439-441; Microscopical Structure, 441-443. Adulteration, 443-445. Added

Oil in Paprika, 445.

Ginger; Composition, 445-446. Exhausted Ginger, and its Detection, 447-448. Microscopical Structure, 449. Adulteration and Standard, 450.

Turmeric; Composition, 450. Microscopical Structure, 451. Detection,

Mustard; Composition, Preparation, 453-456. Mustard Oil Determination, 457. Microscopical Structure, 458. Adulteration and Standards, 459. Coloring Matter, 460. Prepared Mustard; Composition, Adulteration, 460. Analytical Methods, 461.

Nutmeg and Mace; Composition of Nutmeg, 462-463. Microscopical Structure of Nutmeg; Adulteration; Standard of Purity, 464. Composition of Mace, 465. Microscopical Structure; Adulteration; Standard, 466. Bombay or Wild Mace and its Detection, 467. Macassar Mace, 468.

References on Spices, 468.

CHAPTER XIII.

PAGE.

Nature and Properties, 471. Fatty Acids, 471-472. Saponification, 472. Analysis; Rancidity; Judgment as to Purity; Filtering, Weighing, and Measuring Fats, 473. Specific Gravity, 474-476. Viscosity, 477. Meltingpoint, 480. Reichert-Meissl Process for Volatile Fatty Acids, 481-482. Polenske Number, 483. Soluble and Insoluble Fatty Acids, 484-486. Saponification Number, 486. Iodine Absorption Number; Hübl's Method, 487-400. Hanus's Method, 491. Wijs's Method, 492. Bromine Apsorption Number, 402-403. Thermal Tests, 403. Maumené Test, 404. Bromination Test, 404-497. The Acetyl Value, 497-498. The Valenta and Elaïdin Tests, 499. Free Fatty Acids, 500. Titer Test, 500-501. Unsaponifiable Matter, 501. Cholesterol and Phytosterol, 502. Separation and Crystallization, 503-506. Bömer's Phytosterol Acetate Test, 507. Constants of Edible Oils and Fats, 508-509. Parraffin; Microscopical Examinaltion of Oils and Fats, 510. Olive Oil, 511. Composition and Adulteration, 512. Standards, 513. for Adulteration, 513-515. Cottonseed Oil, 516. Bechi's Test, 517. Halphen's Test, 518. Sesame Oil, 518. Adulterants and Tests, 519. Rape Oil, 520. Tests, 521. Corn Oil, 521. Sitosterol, 522. Peanut Oil, 522. Adulterants; Renard's Method, 523. Bellier's Method, 524. Mustard Oil, 525. Poppyseed Oil, 526. Sunflower Oil, 526. Rosin Oil, 527. Cocoanut Oil, 528. Cocoa Butter: Tallow, 520.

Butter, 529. Composition, 530. Effects of Feeding, 531. Analytical Methods, 531. Water, 531-533. Fat, 533. Ash; Casein; Milk Sugar; Lactic Acid; Salt, 534. Standard Butter Fat, 535. Adulteration, 535. Colors, 535-537. Preservatives, 538-539. Renovated or Process Butter, 540. Oleomargarine; Manufacture, 541. Coloring; Detection of Palm Oil, 542. Adulterants; Healthfulness, 543. Distinction from Butter, 544. Distinguishing Tests for Butter, Process Butter, and Oleomargarine, 546. Butyrorefractometer, 546-548. Reichert-Meissl Number; Specific Gravity; Foam Test, 549. Milk Test, 550. Curd Tests, 551. Microscopical Examination, 552-553. Foreign Oils, 554.

Lard, 554. Composition; Lard Oil, 555. Compound Lard; Standards; Adulteration, 556. Foreign Oils, 557. Microscopical Examination, 557-558. Analysis of Lard and Lard Substitutes, 559. Effects of Feeding, 560.

References on Edible Oils and Fats, 561. References on Butter, 562. References on Lard, 563.

CHAPTER XIV.

The Polariscope and Saccharimetry, 578-583. Comparison of Scales and

Normal Weights, 583. Specific Rotary Power; Birotation, 584.

Analysis of Cane Sugar and its Products; Tests for Sucrose, 585. Moisture; Ash; Non-sugars; Sucrose Determination by Polariscope, 586-587. Inversion; Clerget's Formula, 588. Detection and Determination of Invert Sugar, 589. Ultramarine in Sugar; Copper Reduction, 590. Volumetric Fehling Process, 591-592. Gravimetric Fehling Methods, 593. Defren-O'Sullivan Method, 594-597. Munson and Walker Method, 598-607. Allihn Method; Electrolytic Apparatus 608-612. Sucrose Determination by Fehling Solution, 612.

Analysis of Molasses and Syrups, 613. Solids; Ash; Polarization, 613-620. Double Dilution Method of Polarizing; Raffinose Determination, 620. Adulteration of Molasses and Standards, 621. Glucose Determination, 621-624. Ashing Saccharine Products, 624. Tin Determination, 625.

Separation and Determination of Various Sugars, 625–626.

Analysis of Maple Products, 627. Moisture; Ash; Malic Acid Value, 627. Lead Number, 628. Hortvet Number, 628-630. Sy's Method, 630.

Analysis of Glucose; Polarization Formulæ, 630-631. Dextrin; Arsenic in Glucose, 632.

Honey; European, 633. Canadian; American; Hawaiian, 634-635. Adulteration, 636-638.

Analysis of Honey; Moisture; Ash; Polarization, 639. Reducing Sugars; Levulose; Dextrose; Sucrose; Dextrin, 640. Acids; Glucose, 641. Invert Sugar; Distinction of Honeydew from Glucose, 642.

Confectionery; Standard; Adulteration; Colors, 645. Analysis of Confectionery; Mineral Adulterants, 646. Ether Extract; Paraffine, 647. Starch; Polarization, 648. Alcohol; Colors; Arsenic, 649.

References on Sugars, 650.

CHAPTER XV.

Fermented Liquors; Cider, 678. Manufacture and Composition, 678-681. Adulteration, 682. Perry, 683. Wine, 684. Classification of Wines, 685. Composition and Varieties, 686-689. Standards, 689-691. Adulteration, 691-695. Analytical Methods for Wine; Extract; Acidity, 696. Extract Table, 697-699. Tartaric Acid, 701. Malic Acid, 702. Sugars; Glycerin,

703. Tannin, 704. Foreign Colors, 704-706.

Malt Liquors; Beer, 707. Varieties of Beer and Ale, 708. Composition, 709. Malt and Hop Substitutes, 710. Adulteration and Standards, 711. Malted vs. Non-malted Liquors, 712. Preservatives; Arsenic, 713. Temperance Beers, 714. Analytical Methods, 714. Alcohol, 715. Extract, 715–722. Original Gravity, 722–724. Sugars; Dextrin; Glycerine; Acids, 724. Protein; Phosphoric Acid, 725. Carbon Dioxide, 726. Bitter Principles, 726–727. Arsenic, 728. Malt Extract, 729.

AGE

PAGE

Distilled Liquors; Standards for Spirits, 730. Fusel Oil, 731. Whiskey, 731. Manufacture 731-732 Standards, 733-734. Composition, 734-737. Adulteration, 738. Brandy; Manufacture; Composition, 739. Standards, 740. Adulteration, 741. Rum; Composition, 742. Standards, 742-743. Gin, Composition, 744. Analytical Methods for Distilled Liquors; Extract, Acids; Esters; Aldehydes, 745. Furfural, 746. Fusel Oil, 746-749. Methyl Alcohol, 749-752. Caramel, 752-753. Opalescence Test, 753.

Liqueurs and Cordials, 754. Analysis of Liqueurs, 755. References on Alcoholic Beverages; on Beer, 756.

References on Cider and Wine, 757; on Distilled Liquors, 758.

CHAPTER XVI.

Adulteration of Vinegar; Standards, 770-771, Artificial Cider Vinegar, 772. Character of Residue and Ash, 772-773. Character of Sugars, 774. Tests, 775. Composition of Artificial Cider Vinegars, 776. Detection of Adulterants, and Mineral Impurities, 777-778.

References on Vinegar, 778.

CHAPTER XVII. .

Coal-tar Colors, 791. Allowed Colors, 792. Detection in Food; Basic and Acid Dyes; Wool Dyeing, 793. Double Dyeing Method, 794. Vegetable Colors on Wool; Extraction of Colors by Immiscible Solvents, 795. Separation with Ether, 796. Special Tests, 797. Classification and Identification of Coal-tar Dyes; Rota's Scheme, 797-802. Direct Identification of Colors, 803. Table of Reactions for Colors on the Fiber, 804-811. Reagents, 812.

References on Colors, 813.

CHAPTER XVIII.

PAGE

CHAPTER XIX.

CHAPTER XX.

Lemon Extract, 861. Standards, 861. Adulteration, 862. Analytical Methods; Determination of Lemon Oil, 863-865. Alcohol, 866. Citral, 866. 868. Methyl Alcohol; Colors, 869. Solids; Ash; Glycerin; Examination of Lemon Oil, 870. Constants of Lemon and other Essential Oils, 871. Citral, Citronellal, and other Adulterants, 872. Orange Extract, 873.

Almond Extract, 873. Benzaldehyde; Standard, 874. Adulteration; Analytical Methods; Determination of Benzaldehyde, 875. Nitrobenzol; Distinction and Separation from Benzaldehyde, 876. Artificial Benzaldehyde; Alcohol; Hydrocyanic Acid, 877. Wintergreen Extract; Standards; Adulteration; Determination of Wintergreen Oil, 878. Peppermint Extract; Peppermint Oil; Standards, 879. Analytical Methods; Spearmint Extract, 880. Spice Extracts; Standards, 880; Analytical Methods, 881. Rose Extract; Standards, 882. Determination of Rose Oil; Imitation Fruit Flavors, 883. Determination of Esters, 884. Composition of Imitation Essences, 885. References on Flavoring Extracts, 886.

CHAPTER XXI.

PAGE

Canned Vegetables and Fruits, 887. Method of Canning, 888. Composition, and Methods of Proximate Analysis, 889. Decomposition and Detection of Spoiled Cans, 890. Gases from Spoiled Cans, 891. Metallic Impurities, 892. Action of Fruit Acids on Tin Plate, 893–896. Salts of Lead; 896. Salts of Zinc and Copper, 897–898. Greening of Copper Salts, 898. Salts of Nickel; Toxic Effects of Metallic Salts, 899. Separation and Determination of Metallic Salts, 899–903. Antiseptics in Canned Foods, 903. Detection of Preservatives, 904. Soaked Goods, 905. Ketchups and Table Sauces, 905. Composition, 906. Coloring, 907. Preservatives in Table Sauces, 908. Pickles and their Adulteration, 909. Horseradish, 910.

Jams and Jellies, 909. Composition and Adulteration, 911–914. Adulterated Jams and Jellies, 914–915. Labeling "Compound" Goods, 915. Analytical Methods, 916. Determination of Sugars, 917–919. Glucose, 919. Dextrin; Tartaric Acid, 920. Coloring Matters, 921. Preservatives; Starch; Gelatin; Agar Agar, 922. Apple Pulp; Microscopical Examination, 923.

Fruit Juices, 923-924. Grape Juice, 924. Sweet Cider; Lime Juice, 925. Fruit Syrups, 926.

References on Canned Foods and Fruit Products, 927.

PLATES I-XL.

PHOTOMICROGRAPHS OF PURE AND ADULTERATED FOODS AND OF ADUTERANTS

Cereals: Barley, I. Buckwheat, II, III. Corn, III, IV. Oat, IV. V. Rice, V, VI. Rye, VI, VII. Wheat, VIII.

Legumes: Bean, IX. Lentil, IX, X. Pea, X. XI.

Miscellaneous Starches: Potato; Arrowroot; Tapioca, XII. Turmeric; Sago, XIII.

Coffee, XIV, XV. Chicory, XV. XVI. Cocoa, XVI, XVII. Tea, XVIII.

Spices: Allspice, XVIII, XIX. Cassia, Cinnamon, XX-XXII. Cayenne, XXII-XXIV. Cloves; Clove Stems, XXIV-XXVII. Ginger, XXVII-XXIX. Mace, XXIX. Nutmeg, XXX. Mustard, XXXI-XXXIII. Pepper, XXXIII-XXXVI. Spice Adulterants: Olive Stones; Cocoanut Shells, XXXVI. Elm Bark; Sawdust; Pine Wood, XXXVII.

Edible Fats: Pure Butter; Renovated Butter; Olemargarine, XXXVIII. Lard Stearin, XXXIX. Beef Stearin, XL.

FOOD INSPECTION AND ANALYSIS.

CHAPTER I.

FOOD ANALYSIS AND OFFICIAL CONTROL.

INTRODUCTORY.

THE general subject of food analysis, in so far as the public health is concerned, is to be considered from two somewhat different standpoints: first, from the outlook of the government, state, or municipal analyst, whose mission it is to ascertain whether or not the food may properly be considered pure or free from adulteration; and second, from the point of view of the food economist, whose aim is to determine its actual composition and nutritive value. The one protects against fraud and injury, the other furnishes data for the arrangement of dietaries and for an intelligent conception of the rôle which the various nutrients play in the metabolism of matter and energy in the body. The two fields are as a rule distinct each from the other, often involving, in the examination of the food, different methods of procedure.

Official Control of Food.—In view of the importance of the consideration of food with reference to its purity, an ever-increasing number of states have realized the necessity of protecting their citizens from the unscrupulous manufacturers who in various lines are seeking to produce cheaper or inferior articles of food in close imitation of pure goods. Many of the states have laws in accordance with which the sale of such impure or adulterated foods is made a criminal offense, and some, but not all of these, are provided with public analysts and other officers to enforce these laws and punish the offenders. Numerous communities are awake to the importance of municipal control of such commonly used articles of food as milk, butter, and vinegar, and in many cases have machinery of their own for regulating the sale of these foods.

Since January 1, 1907, the federal government has been actively engaged in the enforcement of the national food law of June 30, 1906, through the Bureau of Chemistry of the U. S. Department of Agriculture. In addition to the central laboratories of this Bureau at Washington, upwards of 20 branch laboratories have been established in the principal cities of the United States to enforce the provisions of the national law which regulates interstate commerce in foods, as well as their manufacture and sale in the territories and the District of Columbia, and their importation from foreign countries.

Food Analysis from the Dietetic Standpoint.—The study of the principles of dietetics has been given increased attention during the last decade in the curricula of many of the technical schools and colleges. Much has been accomplished by certain of the state experiment stations working as a rule in connection with the United States Department of Agriculture along this line. Investigations of this character are especially valuable, and are indeed rendered necessary by the general tendency of the modern physician to regard the hygienic treatment of disease, especially with reference to the matter of diet, as often of far greater importance than the mere administering of drugs.

The food economist studies the varying conditions of age, sex, occupation, environment, and health among his fellow men, with a view to showing what foods are best adapted to supply the special requirements of various classes. The quantity and proportion of protein, fat and carbohydrates, or of fuel value best suited for the daily consumption of a given class or individual having been determined, dietaries are made up from various food materials to supply the need with reference as far as possible to the taste and means of the consumer.

Experiments are made on families, clubs, or individuals, representing various typical conditions of life, and extending over a given period, during which records are kept of the available food materials on hand and received during the term of the experiment, as well as of those remaining at the end. In the case of individuals, additional records may be kept of the amount and composition of the urine and feces. From such data the physiological chemist calculates the amount of nutrients utilized, and studies the metabolism of material in the human body.

Up to this point no very extensive apparatus is required, but if in addition the income and outgo of heat and energy are to be studied, which are important to a complete investigation of the economy of food in the body, the student will require a respiration calorimeter and its appurtenances. The calorimeter is so constructed that an individual may be confined therein for a term of days under close observation and with carefully regulated conditions. Such an equipment involves a large expenditure and is to be found in but few laboratories.

It is not the purpose of the present work to go beyond the strictly chemical or physical processes involved in making the analyses by which the proximate components of the foods are determined. For more complete information in the field of dietary studies and the metabolism of matter and energy in the body, the student is referred especially to the investigations of Atwater and his coworkers, as published in the annual reports of the Storrs Experiment Station at Middletown, and in the bulletins of the U. S. Department of Agriculture, Office of Experiment Stations, a list of which is given at the end of Chapter III.

Commercial Food Analysis.—The proper preparation of food products has long ceased to be carried on by the hap-hazard rule-of-thumb methods that formerly prevailed. Now in the manufacture of many prepared foods and condiments, especially on a large scale, it has become a necessity to use scientific processes, rendered possible only by the employment of skilled chemists. In fact it is coming to be more and more common for food manufacturers to establish chemical laboratories in connection with their works, in the interests both of economy and of improved production.

Frequently disputed points arise in the enforcement of the food laws that render the services of the private food analyst of great importance both to manufacturer and dealer. Thus a wide field is open to the analyst of foods outside the domain of the government or state laboratory, either in connection with the large food manufacturing plants directly, or in private laboratories for experimental research, or for analytical control work.

SYSTEMATIC FOOD INSPECTION.

Functions of the Official Analyst.—The public analyst is employed by city, state, or government to pass judgment on various articles of food taken from the open market by purchase or seizure, either by himself or by duly authorized collectors employed for the purpose. The sole object of his examination is to ascertain whether or not such articles of food conform to certain standards of purity fixed in some cases by special law, and in others by common usage or acceptance. Such a public analyst need not concern himself with the dietetic value of the food or whether it is of high or low grade. It is for him to determine simply whether it is genuine or

adulterated within the meaning of the law, and, if adulterated, how and to what extent. Aside from his skill as a chemist, it is often necessary for him to possess other no less important qualifications, chief among which are his ability to testify clearly and concisely in the courts, and to meet at any time the most rigid kind of cross-examination, it being of the utmost importance that he understand thoroughly the nature of evidence.

Standards of Purity for Food Products.*—Under an act of Congress approved March 3, 1903, standards of purity for certain articles of food have been established as official standards for the United States by the Secretary of Agriculture. The earlier of these standards were formulated under the Secretary's direction by a committee of the Association of Official Agricultural Chemists. Later, however, a joint committee of that association and of the Association of State and National Food and Dairy Departments has had charge of this work. Standards have been and are being thus adopted, covering the entire range of food products.

Nature of the Analytical Methods Employed.—Usually but a small percentage of the samples submitted for examination are actually adulterated. The analyst should, therefore, adopt for economy in time the quickest possible reliable processes for separating the pure from the impure, so that most of his attention may be devoted to the latter. The nature of the processes by which this is done varies with the foods. Experience soon enables one to judge much by even the characteristics of taste, appearance, and odor, though such superficial indications should be used with discretion. One or two simple chemical or physical tests may often suffice to establish beyond a doubt the purity of the sample, after which no further attention need be paid to it.

A sample failing to conform to the tests of a genuine food must be carefully examined in detail for impurities or adulterants. While in most cases usage or experience suggests the forms of adulteration peculiar to various foods, the analyst should be on the alert to meet new conditions constantly arising. His methods are largely qualitative, since technically he need only show in most cases the mere presence of a forbidden ingredient, though for the analyst's own satisfaction he had best determine the amount, at least approximately.

In reporting approximate quantitative results in court, especially when they are calculated from assumed or variable factors, or when they are the result of judgment based on the appearance of the food under

^{*} U. S. Dept. of Agric., Off. of Sec., Circ. 19.

the microscope, the analyst should always be conservative in his figures by expressing the lowest or minimum amount of the adulterant, so as to give the defendant the benefit of any doubt. When exact standards are fixed by law, as in the case of total solids or fat in milk, for example, there is of course great necessity for preciseness in quantitative work.

A full analysis of an adulterated food beyond establishing the nature and amount of the adulteration is entirely unnecessary, and in most instances adds nothing to the strength of a contested case, as twenty years' experience in the enforcement of the food laws in Massachusetts has shown.

The responsibility resting upon the analyst is not to be lightly considered, when it is realized that his judgment and findings constitute the basis on which court complaints are made, and the payment of a fine or even the imprisonment of the defendant may be the result of his report. Therefore he should be sure of his ground, knowing that his results are open to question by the defendant. Where court procedure is apt to be involved, a safe rule is for the analyst to consider himself the hardest person to convince that his tests are unquestionable, making every possible confirmatory test to strengthen his position and consulting all available authorities before expressing his opinion; and finally, after being fully convinced that a sample is adulterated, and having so alleged, let him adhere to his statements and not waver in spite of the most rigid cross-examination to which he may be subjected.

While each state or municipality has its own peculiar code of regulations and restrictions concerning the duties of the analyst and other officials, these rules are in the main very similar. For instance, it is usually necessary, excepting in the case of such a perishable food as milk, for the analyst to reserve a portion of a sample before beginning the analysis, which sample, in the event of proving to be adulterated, shall be sealed, so that in case a complaint is made against the vendor, the sealed sample may, on application, be delivered to the defendant or his attorney.

Adulteration of Food.—Except in special cases a food in general is deemed to be adulterated if anything has been mixed with it to reduce or lower its quality or strength; or if anything inferior or cheaper has been substituted wholly or in part therefor; or if any valuable constituent has been abstracted wholly or in part from it; or if it consists wholly or in part of a diseased, decomposed, or putrid animal or vegetable substance; or if by coloring, coating, or otherwise it is made to appear of greater value than it really is; or if it contains any added poisonous

ingredient. These provisions briefly expressed are typical of the general food laws adopted by most states and by the government, though the verbiage may differ. Laws covering compound foods and special foods vary widely with the locality. As to the character of adulteration, nine out of ten adulterated foods are so classed by reason of the addition of cheaper though harmless ingredients added for commercial profit, rather than by the addition of actually poisonous or injurious substances, though occasional instances of the latter are found.

Authentic instances of actual danger to health from the presence of injurious ingredients are extremely rare, so that the question of food adulteration should logically be met largely on the ground of its fraudulent character. Indeed the commoner forms of adulteration are restricted to a comparatively small number of food products, the most staple articles of our food supply, such as sugar and the cereals, eggs, fresh meat, fresh vegetables and fruit being rarely subject to adulteration.

Misbranding.—Under the federal food law and the laws of many of the states misbranding constitutes an offense as well as adulteration. By misbranding is meant any untrue or deceptive statement or design on the label of a food package, either regarding the nature of the contents, or of the place of manufacture or name of manufacturer. One of the commonest forms of misbranding consists in the incorrect statement of weight or measure. Extravagant and untrue claims as to nutritive value have hitherto constituted a frequent form of misbranding.

A Typical System of Food Inspection.—The efficiency of a system of public food inspection is greatly enhanced if the business part of the work, including the bookkeeping and attending to the outside public, be done wholly through some person other than the analyst, as, for example, a health officer, to whom the collectors of samples and the analyst may report independently as to the results of their work, and whose duty it is to determine what shall be done in cases of adulteration. In this way the analyst knows nothing of the data of collection nor the name of the person from whom the sample was purchased, so that he can truthfully state in court that his analysis was unbiased.

Suppose, for example, that three collectors are employed to purchase samples of food for analysis, their duties being to visit at irregular intervals different portions of a state or municipality. Each collector keeps a book in which he enters all data as to the collection of the sample, includ-

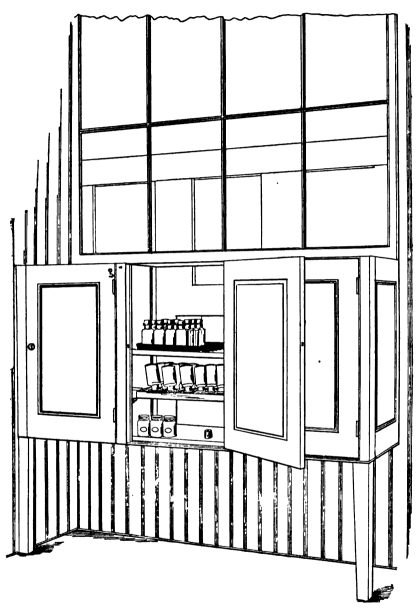


Fig. 1.—Inspectors' Lockers. Insuring safe legal delivery of samples collected by tares inspectors. Each locker has a door in the rear accessible, from an antercom, to the inspector holding key to that locker only.



Fig. 2.—Inspectors' Lockers. Front View. The lockers are accessible to the analyst in the laboratory by a single sliding-sash front, provided with a spring lock. The removable sliding-racks are convenient for returning clean sample bottles.

ing the name of the vendor, assigning a number to each sample, which number is the only distinguishing mark for the analyst. may use for this purpose the odd numbers in succession from I to oooo. the second the even numbers from 2 to 10,000, while the third may use the numbers from 10,000 up. Each of the two former would begin with a lettered series, as, for instance, A, numbering his samples 1A, 3A, 5A, 7A, etc., or 2A, 4A, 6A, etc., till he reached 10,000, then beginning on series B and so on. If the analyst is to be kept in ignorance of the brand or manufacturer in the case of package goods, the collector must remove from the original package sufficient of the sample for the needs of the analyst, and deliver it to the latter in a plain package, bearing simply the name under which the article was sold and the number. Such precautions are, however, not always practicable and depend largely on local regulations. The analyst reports the result of the analysis of each sample with the number thereof on a library card, with appropriate blanks both for data of analysis and for data of collection, the latter to be filled by the collector from his book after the analyst has handed in the card with the data of analysis. This system of recording and reporting analyses has been successfully used for years by the Department of Food and Drug Inspection of the Massachusetts State Board of Health.

Legal Precautions.—The laboratory of the public analyst should preferably be provided with a locker for each collector, to which access may be had only by that collector and the analyst, so that in the absence of the latter, or when circumstances are such that the samples cannot be delivered to him personally, there may be such safeguards with respect to lock and key as to leave no question in the courts as to safe delivery and freedom from accidental tampering. With such a system it is unnecessary for the collector to place under seal the various samples submitted for analysis. Unless such lockers or their equivalent are employed, it is best to carefully seal all samples.

Such a system of lockers for use with three collectors is shown in Figs. 1 and 2. The same careful attention should afterwards be given to keep the specimens in a secure place both before and during the process of analysis, and to label with care all precipitates, filtrates, and solutions having to do with the samples, especially when several processes are being simultaneously conducted, in order that there may be no doubt whatever as to their identity. The importance of precautions of this kind in connection with court work can hardly be too strongly emphasized.

Practical Enforcement of the Food Law.—In the case of foods actually found adulterated, there are three practical methods of suppressing their further sale, viz., by publication, by notification, and by prosecution. These may be separately employed or used in connection with each other, according to the powers conferred by law on the commission, board, or official having in charge the enforcement of the law, and according to the discretion of such official.

Publication.—Under the laws of some states, the only means of protecting the people lies in publishing lists of adulterated foods with their brands and manufacturers' names and addresses in periodical bulletins or reports. Sometimes it is considered best to publish for the information of the public lists of unadulterated brands as well, and, again, it is held that only the offenders should thus be advertised.

Such publication, by keeping the trade informed of the blacklisted brands and manufacturers, certainly has a decidedly beneficial effect in reducing adulteration, and involves less trouble and expense than any other method. It is obviously an advantage, however, in addition to this to be able in certain extreme cases to use more stringent methods when necessary.

Notification and Prosecution.—The adulteration of food is best held in check in localities where under the law cases may be brought in court and are occasionally so brought. The mere power to prosecute is in itself a safeguard, even though that power is not frequently exercised. Under a conservative enforcement of the law, actual prosecution should be made as a last resort. Neither the number of court cases brought by a food commission nor the large ratio of court cases to samples found adulterated are criteria of its good work. Except in extreme cases, it is frequently found far more effective to notify a violator of the law, especially if it is a first offense, giving warning that subsequent infraction will be followed by prosecution. Such a notification frequently serves to stop all further trouble at once and with the minimum of expense. Instances are frequent in Massachusetts where, by such simple notification, widely distributed brands of adulterated foods have been immediately withdrawn from sale.

Massachusetts was the first of all the states to enact pure-food legislation, and for twenty-five years has had a well-established system of inspection, prosecuting cases under its laws through the Food and Drug Department of the State Board of Health. Cases are brought in court with practically no expense for legal services. Complaints are entered by

the collector, or, as he is termed, inspector, who makes complaint not in his official capacity, but as a citizen who under the law has been sold a food found to be adulterated, and who is entitled to conduct his own case, which he does with the aid of the analyst and such other witnesses as he may see fit to employ. Experience is readily acquired by the inspector in conducting such cases in the lower police or municipal courts, where they are first tried, and years ago the services of legal counsel in Massa chusetts were dispensed with as superfluous.*

Statistics in the annual reports of the Massachusetts Board show with what uniform success these trials have been conducted. While more often settled in the lower courts, occasional appeal cases are carried to the superior courts, where the services of the regular district attorney are of course availed of in prosecuting the case.

Such a system as the above, while admirable for a state or city after long experience in the enforcement of food laws in the courts, is obviously impracticable with newly established systems of state food inspection.

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^{*} Where such a practice is in vogue an intelligent inspector must of course be chosen with reference to his ability to do this court work. The food laws are few and simple, as are also the court decisions rendered under them, so that it is no great task for the inspector to become much more familiar with them than the average general lawyer whom he meets in court and who not infrequently consults the inspector for information regarding these laws.

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CHAPTER II.

THE LABORATORY AND ITS EQUIPMENT.

Location.—The selection of a location for a food laboratory cannot always be made solely with reference to its needs and its convenience, but it is more often subject to economic conditions beyond the analyst's control. Under very best conditions, such a laboratory should be situated in a building designed from the start exclusively for chemical or biological and chemical work. Almost any well-lighted rooms in such a building can be readily adapted for the purpose. When, however, as is frequently the case, rooms for such a laboratory are provided in municipal, government, or office buildings, in which for the most part clerical work is done, the problem of adequately utilizing such rooms so that they may not at the same time prove offensive to or interfere with the comfort of other occupants of the building is sometimes difficult. It is obvious that basement rooms in such a building, as far as ventilation is concerned, are less readily adapted for the requirements in hand than are those of the top floor, though, if the light is good and there are abundant and well-arranged ventilating-shafts, such rooms may be made to serve every purpose. In the basement one may most easily obtain water, gas, and steam, and dispose of wastes without annoyance to one's neighbors. When, however, it is possible to do so, rooms on the top floor of an office building should be utilized for a food laboratory, for in such rooms the problems of lighting, heating and ventilating are comparatively simple and may usually be solved without regard to other occupants. In such a case ample provision must be made, preferably through shafts which are readily accessible for water-, gas-, steam-, and soil-pipes passing down below.

The actual equipment of the food laboratory depends of course largely on its particular purpose; and while it is manifestly impossible to do otherwise than leave the details to the individual taste and needs of the analyst,

modified by the means at his disposal, a few general suggestions regarding important essentials may prove helpful. These imply a fairly liberal though not extravagant outlay, with a view to saving both time and energy by convenient surroundings well adapted to the work in hand. At the same time equally satisfactory work is possible under simpler conditions than those described.

Floor.—The best material for the floor of the working laboratory is asphalt. Such a floor is firm but elastic, is readily washed by direct application of running water, if necessary, and resists well the action of ordinary reagents. An occasional thin coating of shellac with lampblack applied with a brush gives the asphalt floor a smooth, hard surface and may be applied locally to cover spots and blemishes.

Lighting.—The lighting of the rooms, if on the top floor, is best effected by both wall windows and skylights. North windows furnish the best light for the microscope; the skylight, when available, is the ideal light for the balance and for general laboratory work.

Ventilation by forced draft is a great convenience. For this purpose an exhaust-fan driven by an electric motor and controlled in speed by a fractional rheostat is admirable. Such a fan had best be located in a small closed compartment or closet near the centre of the series of rooms designed to be ventilated by it, and this closet should have directly over the fan an outlet-shaft passing through the roof of the building. With such a system, a series of branching air-ducts should radiate from the fan closet, conveniently arranged either above or along the ceiling and communicating with the various hoods, closets, and rooms near the top.

Benches.—The working benches should have wooden or glazed tile tops. White glazed tile, if properly laid, furnish a very clean, sanitary, and resistant surface, besides being often convenient for color tests. If laid on a plank surface, cement should not be applied directly, as it swells the wood before drying out and results in a loose and often uneven surface. Cement may be avoided altogether and the tiles after first soaking in oil may be laid in putty directly on the wood. Tiles may be laid in cement by first covering the plank surface with cheap tin plate, overlapping the edges and securing by tacks. This prevents swelling of the wood. The tin may be covered to advantage with cheap paint. The tiles may then be embedded in a layer of cement spread over the tin surface.

Soft encaustic glazed tiles commonly used for wall finish are not as

effective as hard floor tiles, since the former crackle and lose color when subjected to heat. If the hard floor tiles can be specially glazed, they make by far the most satisfactory and enduring surface.

When wooden bench tops are used they may be treated to advantage by staining with the following solutions:

Solution 1. 100 grams of anilin hydrochloride, 40 grams of ammonium chloride, 650 grams of water.

Solution 2. 100 grams of copper sulphate, 50 grams of potassium chlorate, 615 grams of water.

Apply solution I thoroughly to the bare wood and allow it to dry; then apply 2 and dry. Repeat these applications several times. Wash with plenty of hot soap solution, let dry and rub well with vaseline. It is claimed that wood so treated is rendered fire-proof and is not acted on by acids and alkalies. When the finish begins to wear, an application of hot soap solution or vaseline will bring back the deep black color.

The benches should naturally be located with reference to best light from skylights or windows. Gas and water outlets, sinks and waste-pipes should be conveniently arranged with reference to the working benches, as well as suitable provisions for air-blast and exhaust, while in the space beneath the benches such drawers, cupboards, and receptacles as are required should be provided. A clear bench width of 24 inches is ample for most work; if wider there is a temptation to allow apparatus to accumulate at the back. At the back of the bench and within easy reach, a raised narrow shelf should be provided to be used exclusively for common desk reagents. This again should not be so wide as to allow the accumulation of useless bottles. A narrow raised guard or beading at the edge of the reagent shelf prevents the bottles from accidently slipping off.

Hoods.—Closed hoods with sliding sash fronts are almost indispensable. These hoods should be directly connected with the ventilating shafts or pipes, or with the air-ducts that radiate from the exhaust-fan closet, when such a system is provided. Gas outlets inside the hoods are necessary.

When there is a good draft, either natural or forced, a hooded top over the working bench, such as that shown in Fig. 3, is quite as efficient as a closed hood for most purposes. This is best made of galvanized iron, painted on the outside and treated on the inside with a preparation of graphite ground in oil. Here are best carried out all the processes involving the giving off of fumes and gases, which, if the ventilation is efficient, should pass directly up the flues and not come out in the room,

Sinks and Drains.—The sinks should preferably be of iron or porcelain. If iron, they should at frequent intervals be treated with a coat of

Fro. 3.—Hooded Top of Galvanized Iron over Working-bench, Connected with Ventilating Air-ducts.

asphalt varnish. A great convenience is a hooded sink (Fig. 4) in which foul-smelling bottles, or vessels giving off noxious or offensive fumes

or gases, may be rinsed under the tap while completely closed in. Openwork rubber mats at the bottom of the sinks help to insure against breakage. Open plumbing of simplest design should be used, and a multiplicity of traps should be avoided. Sinks may be variously located for

F10. 4.—A Hooded Sink. An injector-like arrangement of steam and cold-water pipes furnishes water of any desired temperature.

convenience without regard to situation of soil-pipes, if the floor is thick enough to allow an open drain with sufficient pitch to flow readily. Such open drains are much more readily cleaned than closed pipes, and are best constructed by splitting a lead pipe and laying it in an iron box which is sunk into the floor. The edges of the lead pipe are rounded over those of the box as in Fig. 5, filling the joints with hydraulic cement, and the top of the drain is covered by a series of readily removable iron plates

flush with the top of the floor. Waste-pipes from sinks, still-condensers, refrigerators, and various forms of apparatus involving flowing water may be led into this drain, holes being drilled in the iron cover for their insertion.

Steam and Electricity.—These are useful but not indispensable. Steam, when available, may be used to advantage for boiling ether or benzine in connection with continuous fat-extraction apparatus, for furnishing the motive power for driving the Babcock centrifuge, for heating waterbaths and hot closets, and, in connection with cold water, to furnish a

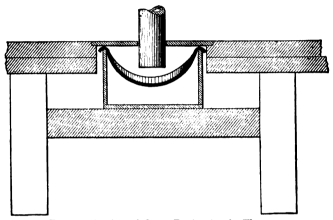


Fig. 5.—Section of Open Drain-pipe in Floor.

supply of hot water when wanted at the sink. The latter application is illustrated in Fig. 4.

If electricity is used for lighting, it may also be applied in a variety of useful ways in the laboratory, as, for instance, for heating coils or electric stoves, for electrolysis, and for running small motors, which in turn may be employed for driving centrifuges, shaking apparatus, ventilating-fans, air-pumps, etc.

Suction and Blast.—If the water-pressure is ample, both air-pressure and exhaust for blast-lamps, vacuum filtration, and other purposes are readily available through the agency of the various devices used in connection with the flow of water, as, for instance, the Richards pump. When, however, the water pressure is insufficient, other means must be employed for furnishing these much-needed requisites. Fig. 6 illustrates a simple and almost noiseless pressure and exhaust pump run by a $\frac{1}{8}$ -H.P. electric motor, which with the pressure-equalizing tank and the appropriate connections are mounted on a light wheel truck, and readily movable to any part of the laboratory. By simply screwing the plug into an

electric-light outlet, either suction or blast may be had at will, depending on the position of a knife-edge switch which determines the direction of the current. By means of a fractional rheostat the speed may be varied and the pressure thus controlled.

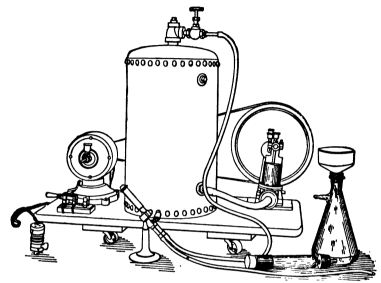


Fig. 6.—Portable Pressure- and Exhaust-pump Run by Electric Motor. Useful for blast-lamps, vacuum filtration, etc.

APPARATUS.

The laboratory is of course to be supplied with the usual assortment of test-tubes, flasks, beakers, evaporating and other dishes of porcelain, platinum and glass, funnels, casseroles, crucibles, mortars, burettes, pipettes, graduates, rubber and glass tubing, lamps, ring-stands and various supports, clamps and holders, the nature, number, and sizes of which are determined by individual requirements. Special forms of apparatus peculiar to certain processes of analysis or to the examination of special foods will be described in their appropriate connection. The following apparatus of a general nature may be regarded as indispensable for the proper fitting out of the food laboratory:

Balances.—These should include (1) an open pan balance for coarse weighing, having a capacity up to 1 kilogram and sensitive to 0.1 gram, with a set of weights; and (2) an analytical balance, enclosed in a case, sensitive to .0001 gram under a load of 100 grams, with an accurate set of non-corrosive weights. The short-beam analytical balance is prefer-

able for quick work, and as constructed by the best modern makers leaves nothing to be desired.

The Water-bath.—This is such an important accessory to the food analyst that it should, if possible, be specially designed to meet his require-

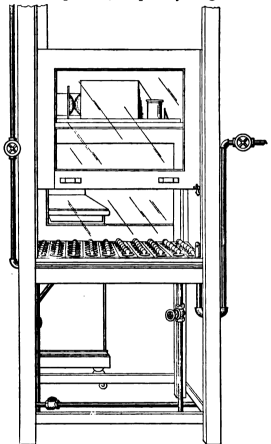


Fig. 7.—Water-bath, Enclosed in Hood, with Sliding-sash Front.

ments, though the ordinary copper baths, supported on legs and designed to be heated by gas-burners, as kept in regular stock by the dealers, will sometimes serve the purpose. For nearly all moisture determinations the platinum dishes described on page 133 and the somewhat larger wine-shells of 100 cc. capacity are most used, and for this purpose the top of the bath should have plenty of openings of the right size for these. A very economical construction of bath admirably adapted for the food analyst's use is shown in plan in Fig. 8, being the form employed by the writer.

The size and number of openings are determined by the number of samples to be simultaneously analyzed. The dotted lines indicate a steam-coil within the body of the bath, which serves to boil the water. Fig. 7 shows the bath in place within a hood, the sliding front of which is furnished with a hasp and padlock, so that it may always be kept locked by the analyst whenever he is temporarily absent from the laboratory. This is a useful precaution, when the residues left thereon are from samples which are to form subjects for possible prosecution in court later.

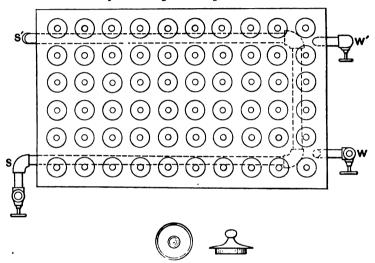


Fig. 8.—Plan View of Water-bath Boiled by Steam-coil. W and W are water inlet and outlet, S and S steam inlet and outlet, respectively.

Steam, if available at all seasons of the year, furnishes a ready means of heating the bath. Electric immersion coils are also convenient. In the absence of both steam and electricity, the bath must be boiled by gasburners.

The Drying-oven.—A convenient form of asbestos-covered, jacketed air-oven, having removable shelves and heated by a gas-burner is shown in Fig. 9, with an efficient form of gas-pressure regulator. The particular form of low burner best adapted for use with the oven is also shown. Such an oven may also be heated by an electric coil, the temperature being governed by a rheostat of delicate construction.

The Water-still.—An efficient still should be provided, capable of supplying the laboratory with an ample quantity of pure water for analytical purposes. Fig. 10 illustrates a compact form of still, which is particularly economical in view of the fact that a single stream of inflowing cold

water first serves to cool the condenser, and, rising, becomes vaporized in the boiler directly connected with the condenser at the top. This apparatus is capable of distilling six gallons of water in twelve hours.

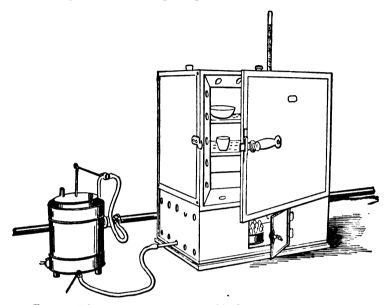


Fig. 9.—Asbestos-covered Air-oven, with Gas-pressure Regulator.

The list of indispensable requisites in addition to the above should include the following:

Continuous Extraction Apparatus (Figs. 18, 19, and 20).

Apparatus for Nitrogen Determination (Figs. 24, 25, and 26).

Apparatus for Distilling Various Food Products (pp. 71 and 660).

A Babcock or other Milk-fat Centrifuge (Figs. 11, 44, and 45).

A Butyro Refractometer (Fig. 36).

An Immersion Refractometer (Fig. 40).

A Microscope and its Appurtenances (Chapter V).

A Polariscope and its Accessories (Figs. 102, 103, and 104).

Apparatus for Specific Gravity Determination (Figs. 14, 15, 16 and 17).

Apparatus for the Determination of Carbon Dioxide (Fig. 71).

Apparatus for the Determination of Melting-points (Fig. 93).

Marsh Arsenic Apparatus (Fig. 27).

Electrolytic Apparatus (Fig. 110).

Separatory Funnels (Figs. 22 and 23).

Following is a list of apparatus and appliances which, while not indispensable, are convenient and at times desirable:

A Spectroscope, either of the direct-vision variety for the pocket, or the Kirschoff & Bunsen style on a stand.

Spectroscope Cells, parallel-sided, for observation of absorption spectra.

A Photomicrographic Camera and Appurtenances* (pp. 96 to 98).

A Muffle Furnace.

F10. 10.—A Convenient Laboratory Water-still with Earthenware Receptacle, Provided with Faucet and Glass Gauge.

An Incinerator for a Large Number of Residues (Fig. 53).

An Ebullioscope (Fig. 113).

An Assay Balance, for weighing arsenic mirrors to 0.01 mg.

An Abbé Refractometer (Fig. 39).

A Schreiner Colorimeter (Fig. 28).

A Lovibond Tintometer (p. 78).

^{*} A photographic dark room is also necessary if photomicrographic work is to be done-

A Universal Centrifuge.—This convenient apparatus merits a separate brief description, being useful for a wide variety of purposes, such as breaking up ether- and other emulsions, quickly settling out precipitates, and roughly estimating chlorides, sulphates, phosphates, etc., by the volume of the precipitate in graduated tubes. Various-sized aluminum frames, carrying hinged shields, are interchangeably adjustable to the

Fig. 11.—The Universal Centrifuge. Driven by an electric motor.

spindle of a vertical electric motor.* The smallest frame has shields adapted to hold two graduated glass tubes of 15 cc. capacity (see Fig. 11). This is for the quantitative estimation of small precipitates and the quick settling of sediments. A medium-sized and large frame carry tubes of 80 cc. and 120 cc. capacity respectively. A frame is also provided with shields adapted for various-sized beakers to be used in settling precipitates. The milk-fat centrifage frame shown in Fig. 45 is also adapted to be used on the spindle of the same motor.

^{*} In the absence of electricity a water-motor may be used.

FOOD INSPECTION AND ANALYSIS.

REAGENTS.

				-																			
Remarks, Preparation, etc.	Sp. gr. 1.080. '' ' 0.833 to 0.895. B.P. about 70° C. '' '' 1.055.		Sp. gr. 1.20. 38.92% HCl.	r vol. No. 8 to 8 vols. H2O.	Sp. gr. 1.20.	11.42. 69.2% HNOs.		Weigh freshly crystallized No. 16.	Sp. 81. 1.725.	so to 100 Dissolved in 2.5% HCl. See also No. 197.	Saturated.		So or 18436 100% H.SO.	1.185. 25.27% HaSO. 1 vol. No. 27 to 5 vols. HaO.	For Keichert number. For crude fiber.	Standardize against No. 241.	The fact that th	39. 87. 0.516. D.F. 137 C			0.9368. 105 cc. No. 36	0.814.	In alcohol, see page 92.
Grams per Liter.		Ş	704			983 386.5		6.3		50 to 100		100	1843	200	1 20	4			:			:	92
Molec- ular Weight.	102 88 60	62 4	36.4			63	961		2630	229	126	394	138	3 :			322	8 9	:			33	904.4
Designation.	Glacial. 99.5%	Crystallized	C. P. concentrated	10% solution	Commercial, conc.	C. P. concentrated Dilute	Commercial conc	N/10 solution	Crystalliz.d	Solution	Solution Crystallized	10% solution	Crystallized	Dilute	10%01	N/10	Dry	Absolute	94% by vol. U. S. P	50%	25%	Dry root.	Grystallized
Formula.	(CH ₃ CO),O CH ₃ CO,(C ₃ H ₃) H.C ₂ H ₃ O ₂	н,во, с,н,он	нсі			HNO3	C.H. C.H. T.H. C.J.		H11PW10038,8H20	C,H ₂ (NO ₂),OH.	C,H ₃ (OH) ₃ .	C20H16O3	Си, он сол.	ngo(4			C4H10O	CH10H CH10H				сн. он.	Alg(SO4)3-(NH4/35O4+
Name.	Acetic anhydride. Acetic ether. Acid, acetic.	boric.	hydrochloric	::		nitric		OXE	phosphoricphosphotungstic		pyrogallic	rosolic	salicylic	embumuc	::		tannic	Alcohol, amyl.				Alkanna methyl	imonia)
Š	H 4 10 4	100	~so 0	2 :	12	13	1 2	12	8 6	2 2	2 2	7 7	9	287	9	319	33	4 %	300	387	8 9		

	100 grams MoO, No. 185 mixed with 400 cc. cold H ₂ O and 80 cc. ammonia No. 47. Filter and add filtrate tu 300 cc. HNO, No. 13 and 700 cc. H ₂ O.	Saturated. Saturate 3 vols. No. 48 with H ₂ S and add 2 vols. No. 48. See page 181. Pulvenzed bone charcoal, page 614. Washed in acid and alkali, page 594.	Dissolve 4.95 grams As ₂ O ₂ No. 66 and 20 grams K ₂ CO ₃ in 250 cc. H ₂ O. Dilute to 1 liter. 1 gram surcess in ro cc. HCI No. 8. 14 grams crystallized copper acetate No. 104 and 5 cc. acetic acid No. 4 in 200 cc. H ₂ O.	2 grams AgNO ₃ in 200 cc. alcohol. No. 36, and 40 cc. ether. Add drop HNO ₃ .	See perincipalm enterings. 1907-1918. 4 grams bismuth subtrate, 4 grams Rochelle salt, 8 grams sodium hydroxide dissolved in 100 cc. water by aid of heat. See chlorinated lime. See the water because of the salt bromine having a few drops bromine.	in excess in reagent bottle. See page 236.
Crystallized 28% Concentrated 28% Dilute 10% NH+ Solution 114 Solution 233 Dry salt 53.38	Counties 80 Crystallized 142 Solution Crystallized 132	<u> </u>	substance	Lry salt Crystallized 244 Solution 261 Solution 75	Crystalline powder. Crystallized	Crystallized 219
Al ₄ (SO ₄) ₂ K ₅ SO ₄ + p ₀ 24H ₂ O NH ₄ OH (NH ₄) ₂ CO ₅ H ₂ O. NH ₄ CI.	molybdate. (NH4,2MoO4. mitrate NH4,NO5, H2O ozalate (NH4,2MPO4. (NH4,2MPO4. NH4,MO4.	(NH,)S NH,CNS C,H,NH,	Asych	Ba(C ₂ , 2H ₂ O Ba(NO ₃) ₂	BiO.NO ₃ + H ₂ O. Na ₃ B ₄ O ₇ 10 H ₂ O. Br.	CaCl ₂ , 6H ₂ O.
potash) una cream'' un hydrate nia nium carbonate chloride	: :::::	so sulphide sulphide sulphide sulphide sulphocyanate. A Anime charcoal state subsector fiber subsector fiber sulphide su	Badouin's reagent. Barloed's copper acctete reag't.	Darrum Carlonnace. chloride intrate Bechi's reagent.	76 Bismuth solution (alkaline). 77 Bismuth submitrate 80 Beaching powder. 78 Borax. 79 Bromine.	Brücke's reagent. Calcium chloride.

REAGENTS-(Continued).

	For absorbing moisture. See chlorinated lime, No. 95. A saturated solution. A solution in xylol. See sucrose. (1) Heat MnOr with HCl. (2) Treat chlorinated lime	Calcuum hydrate saturated with Cl. H ₂ O saturated with Cl. See page 91. Indicator. Macerate in 25% alcohol. Filter, and neu-	See page 93. See page 93. See pages 322, 339. See pages 422, 339. See pages 422, 339. See pages 422, 339.	See page 66. See page 66. See No. 1. 34.619 grams pure CuSO, 5 HyO in soo cc. HyO. 173 gras. Rochelle salts and so gras. NaOH in soo cc. H ₂ O.
-	92	, o N		100
eger Ben an	4 0 6	290.11	2 663 5 146 63 5 146 63 5	466
	Solution. Anhydrous Granular Slaked lime Oucklime. Dry salt Solution. Liquid. Crystallized Solution. Gas	Crystals Solution Dry. Solution.	Crystalized. Crystallized. U. S. P.	Absolute. Dry salt Solution. Crystallized.
TOTAL STREET	Ca(OH); CaSO, *H40 CaSO, *H40 CS; CCC, CAHCLO.H40	CHCla.	Cu(C ₂ H ₂ O ₂) ₂ H ₂ O. CuSO ₄ sH ₂ O. C ₄ H ₄ C ₂ H ₂ CO ₃ . H ₂ O. C ₂ H ₄ B ₃ CO ₂ . (C ₂ H ₂ B ₃ O ₂ .	Fe ₂ (C ₂ H ₂ O ₂) ₆ . Fe ₂ Cl _{6-1,2} H ₂ O. Fe ₂ SO ₄₋₇ H ₃ O.
	Calcinum chloride. hydrate hypochlorite. oxide. sulphate Canada balsam. Carbon bisulphide. Carbon bisulphide. Carbon tetrachloride Carbon tetrachloride Chlorial hydrate.	Chlorinated lime. Chlorine water. Chloroform. Chlor zinc iodide. Cobalt nitrate.	Collodion silk., Copper acetate Sulphate. Wire Coumarin Cuprammonia Distilled water. Eosin.	Ether Ethyl acetate. Fehing copper solution. Ferric acetate. chloride.
	80 80 80 80 80 80 80 80 80 80 80 80 80 8	200000000000000000000000000000000000000	00000000111	2 420 200 00

Prepare fresh for use. 40% known as formatin.	See page 370. Microscopic mountant, see page 86. Equal vols. amyl alcohol and CS ₂ (containing 1% subprue page 491.	Conc. commercial HCl, with 2.5 cc. No. 17 Treat FeS with dilute H ₂ SO ₄ . Saturate H ₂ O with H ₂ S. See page 487. See page 370. See page 92.	Triturate I in mortar with 18 grams KI and small portions H ₂ O till all is desolved. Dilute to 1 liter, standardizing against No. 245. 2 grams crystallized KI dissolved in 100 cc. H ₂ O. Saturate solution with I. Of known purity.	In 50% alcohol. For indicator. See page 856. Or prepare as on page 586. See petroleum ether. Saturate H ₂ O with Ca(OH) ₃ . Digest 1 page 850.	Dip stripe In alcohol.
88			13.06	*	8
80	0.	44 44 44 44 44 44 44 44 44 44 44 44 44	8 · · · · · · · · · · · · · · · · · · ·	0.5	84
Solution. Solution Crystallised.	Solution U. S. P.	Gas Solution Crystals Resublimed	N/ro solution Solution Powdered	Inty powder. Inducator solution. Dry powder. Crystallized. Solution. Use U. S. P. solution. Dry color. Tincture.	Dry color. Solution. Crystafilised
Pes. CoHishsHCi. CaHsO.COH.	Çah,(OH),	N. N	10th	Carradii H.O. Pb(Carradii H.O.	MgCO, MgCL, 6H,0.
Ferrous sulphate. Rormaldehyde. Puchain. Fuller's earth.	Glycerin jelly Glycerin jelly Halphen a reagent.	Hubl's iodine reagent Indigo carmine.	in potassium lodide. trichloride. Kaolin.		Litmus paper, Logwood, Magnesium carbonate chloride.
	-00 00 = 6 : 6 6 6 7 7 7 1			*****************	159 160 161 161 163 163

REAGENTS-(Continued).

Remarks, Preparation, etc.	Dissolve 100 grams MgCls in HO. Add NH,OH in excess and NH,Cl till precipitate is redissolved. Di-	lufe to 1 lifer.	Pages 233, 284 13.525 grams HgCl ₂ and 40.62 grams KI in 1 liter of water.	Saturate H ₂ O with HgCl ₂ .	See page 147, sugar clarifier.	go% ethyl alcohol No. 36 and 10% methyl alcohol No.41.	In H.O. Free No or formaldehyde.	Sec No. 149. Dissolve Hg in twice its weight of HNOs No. 13 and childre with equal vol. of Ho.	See Nos. 190-192.	Dissolve to grams KI in H ₂ O, add saturated HgCl ₂ solution No. 17, an a distinct red prepit tate. Add 350 ec. of a solution of KOH containing 500 grams per liter, make up to a liter and allow to settle, using	Control of the contro	B.P. 3.ºº to 50° C. for Soxhlet extraction. B.P. 9.º C. and over for cleansing. See No. 6.	See page 90. Dissolve in 50% alcohol.
Grams Per Liter.				. 02									
Molec- ular Weight.	40.3	246.3	: : 6	272	200				143.8	:			31.00
Designation.		Crystallized		Saturated solution	Dry salt		Indicator solution		Dry substance				Dry powder Indicator solution
Pormula.	MgO.	MgSO ₄₋₇ H ₂ O MnO ₂ .	5	HgC1	HgO. HgSO.				MoO ₂ Dry subst				C ₂₀ H ₁₄ O ₄ .
Name.	Magnesium oxide	23		, ig	oxide sulphate. Mercury, acid nitrate of	Methylated spirit.	Wilk.	-sugar.	Molybdic oxide. MoOg. MoDp. Naphtha. Naphtha.		Palas reagent. Paraffin. Petroleum ether.		hydrochlone acid. CasH ₄ O ₄
, o	163	165			175	22.081	1832	184	28 3	100	188	101	105

Dissolve 120 grams sodium phosphate and 200 of sodium tungstate in 1 liter of H ₂ O and add 100 cc. of conc. H ₃ O ₄ . See also No. 20. Properly hydrochloroplatinic acid.	Dissolve in H ₂ O and dilute to 1 liter.		,	See page 181	riepara ireany tor use.		For CO ₂ determination.		Adjust strength of solution by repeated trial, standardising against No. 145.			Dissolve 1 gram of inchisin in H ₂ O add a mixture of 20 cc. NaHSO ₄ solution No. 234 and 10 cc. HCl No. 8. Make up to 1 liter.	See page 93.	Standardize against No. 238.	. Keep under petroleum. See Rochelle salt. No. 224.	Saturated			
	° 4		8	£ .	χ ς,	005	1000	100	3.16		100		: 3	17				150	6.60
	205	138	\$ 50	329	422	26	991	101	3 :	174 174 110	282		170			104	901	58.5	•
Crystallized.	Solution. Dry salt. N/10 solution.	Lystallized salt	Solution Dry salt.	Solution Dry salt		Dry substance.	Crystallized	Solution. Crystallized.			Solution		Crystallized.		Metal	Crystallized.	Dry substance.	Solution.	Dry substance
H ₂ PtCL6H ₂ O	K,Cr,O,	KKIOS KCIOS KCIOS	KCN	KaFe(CN)6	K, Fe(CN) 6.3H2O.	кон	KI	KNO. KMnO.		K.SO. C.H.(OH)2	KNaC,H,O. 4H2O				Z.B.	NaHSO, H2O	Na ₂ CO ₃	NaCL	NaOH
Phosphotungstic reagent	Potassium bichromate.	Fotashum disuphate			ferrocyanide	hydroxide	iodide	nitrate	:	phosphate	Rochelle salt.	Schiff's reagent.			Sodium and potassium tartrate	bisulphite	borate		hydroxide
		0 0 0 0	200	200	2 2 2 2	213	215	218	220	222				3 6 7		233	235	230	230

REAGENTS-(Continued).

Grams per Per Liter.	₹	Agitate 1. with 10 parts H.O.	Ciprace supence, and sphon on supernatant liquid.	Saturate conc. HCl with tin, dilute with an equal vol. of H ₂ O, and from time to time add slight excess of acid. Keep pieces of tin in reagent bottle.	H.O. Pour into too times Boil, settle, and use super-	See actd, tannic, No. 33. See page 519. Make strong alcoholic solution of No ssq, dip strips of filter-paper therein, and dry. See page 725, 778. See page 530. See page 492. As a medium for Canada baheam. See No. 98. A saturated aqueous solution of the salt No. s65 acidibided with HySO.
Molec- G	<u> </u>	:	60 60 - H	<u>: </u>	:	
	Solution.	Solution			Indicator solution	Crystallised Granulated Dry powder. Crystallized Solution Crystallized Solution Crystallized Solution Granulated.
			Na, HPO, 112H-0. Na, S, O, 5H-0. Na, WO, H-0.	SnCl ₂ , rH ₂ U		C ₁₂ H ₂₂ O ₁₁ . Sn. Sn. (UO ₂)(C ₂ H ₂ O ₂) ₂ zH ₂ O. C ₄ H ₁₀ . Zn(C ₂ H ₃ O ₂) ₂ zH ₂ O. ZnSO ₄ zH ₂ O.
	Sodium hydroxide	" hypochlorite	phosphate. thiosulphate	Stannous chloride.	States allowings.	Sucrose. Subhindigotata. Tannin The Tocher's reagent. Turmeric paper. Uranium acetate. Vanilin. Villevechia reagent. Xylul. Zinc. chloriodide. chloriodide.
\	0 17	343	244		0	0 = 1 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4

REAGENTS-SUPPLEMENTARY LIST.

Remarks, Preparation, etc.	Sp. gr. 0.790 B P. 22°-C. See page 746. Dissolve o.s gram in 150 cc. of 20% acetic acid See page 322. Saturated solution.	s parts amyl acetate + 1 part amyl valerianate. 5 cc. C. P. amilin, shaken with 3 cc. water and cleared with 2 cc glacial acetic acid. See page 642.	Dissolve in conc. H ₅ SO ₄ . See page 168. Dissolve 173 grams cryst. Rochelle salts and 125 grams make up to 500 cc. re page 746.
Grams per Liter.	8		10
Molec- ular Weight.	209.1 957.4 957.4	178 178 178 178 185 185 185 185 185 185 185 185 185 18	109
Designation.	Absolute Standard solution Liquid Crystallised Solution Crystallized Solution Crystallized Solution Crystallized Solution	Liquid Solution Crystallized Solution Crystallized Crystallized Crystallized	
Pormula.	CH, CHO. CH, CO CH, NH, C, H, SO, H + 2 H, O (NH,)2 Fee (SO,), + 24 H, O	CaHii CaHiO; CaHii CaHiO; CaHin H CaHiO; + Aq. Ba(CaHiO;) + 2HiO Ba(OH); + 8HiO Cu(NO;); + 3HiO	CuSO ₄ (CaHa) ₂ NH
Name	Acetic aldehyde Acetone Acid, sulphanilic Alum, ferric Ammonium acetate	Amyl acetate valerianate reagent Andin acetate Barnum acetate hydroxide Copper nitrate	Diphenylamin Allihn alkali solution
No.	40 00 HE 40		

REAGENTS-SUPPLEMENTARY LIST-(Continued).

	Name.	Formula.	Designation.	Molec- Grams ular per Weight. Liter.	Grams per Liter.	Remarks, Preparation, etc.
289 G	289 Glycerin-soda	ide	2% solution			20 cc. 50% solution sodium hydroxide + 180 cc. glycerin. See page 482. S
N	Naphtylamine hydrochloride (alpha)	ydrocalonde CigHtNHa.HCl Crystallized 179.5 Solution	Crystallized	179.5		Dissolve o.2 gram in 20 cc. conc. acetic acid with heat, decent, and make up to 150 cc. with 20% acetic acid. See page 322.
2 2 2 2 2 2 4 2 2 2 2 2 2 2 2 2 2 2 2 2	Pancreatin Phloroglucinol Parachen vienediamin	Powdered Powdered Solution C4Hs(OH)++2HsO Crystallized COHSIN CAH (NHs)+ CAH (NH	Powdered Solution Crystallized Crystallized		•	Water solution. Por purification see page 287.
	Potassium hydroxide.	Solution Solution Alcoholic solution	Solution Alcoholic solution		% Q	Dissolve to grams potassium hydroxide free from car-
300	299 300	nganateStandard solution	Standard solution Solution		2 8	See page 749.
301 Pr	ride (meta)	1 nygrocnio- Cala(NHs)sHCl Crystallized calaborer co. 8	Crystallized	. 0	:	For making aldehyde-free alcohol, see page 745.
304 SSI 305 SSI 305 SSI 305 SSI	303 Silver nitrite 304 Sodium chloride 305 Zinc nitrite	AgNO9 Crystallized 153.9 Saturated sol NaNO9 Standard solution See page 322. Dust	Crystallized Solution Standard solution Dust	153.9		Saturated solution. See page 322.

REAGENTS.

The foregoing list includes the general reagents used in carrying out the processes treated of in this volume, together with their strength, mode of preparation when necessary, and other data.

Reagents, especially those constantly employed, should be assigned to regular places on the shelves, and should invariably be kept in place when not in use.

Among the standard solutions for volumetric work, none is more frequently of service in the food laboratory than a tenth-normal solution of sodium hydroxide, and a large supply of this reagent, carefully standardized, should be at all times conveniently at Besides being useful for standardizing tenthnormal solutions, it is constantly needed for determining various acids in food products, such as milk, vinegar, butter, lime juice, cream of tartar, liquors, and many others. Time is well spent in carefully adjusting this solution to its exact tenth-normal value, thus simplifying the calculation of results. stock bottle (say of two gallons capacity) containing the standard tenth-normal sodium hydroxide, is conveniently mounted with a side-tube burette in connection, in some such manner as shown in Fig. 12. A small connecting side bottle contains a strong solution of sodium hydroxide (reagent No. 240) through which

FIG. 12.—Stock Bottle of Tenth-normal Alkali.

the air that enters the large bottle is passed, thus depriving it of CO₂. In this manner the standard solution may readily be kept of unvarying strength for a year or more.

EQUIVALENTS OF STANDARD SOLUTIONS.

No. 31.	DECINORMAL SULPHURIC ACID. C	ne cc. is equivalent to	
	Ammonia gas	NH ₃	.0017 gram.
	Ammonia	NH,OH	.0035 "
	Ammonium carbonate	(NH ₄) ₂ CO ₃	.0046 "
		(NH ₄) ₂ CO ₂ H ₂ O	
		CaCO,	••
		Ca(OH)	•
	•	CaO	0,
		Pb(C ₂ H ₂ O ₂) ₂ , 3H ₂ O	
		MgO.	•
	· ·	MgCO ₃	
		N	
		KC ₂ H ₃ O ₂	
		KHCO ₃	
	Ditartrate	KHC ₄ H ₄ O ₄	
	carbonate	K,CO,	
		$K_2C_0H_5O_7H_2O$	
	nydroxide	КОН	•
		KNaC ₄ H ₄ O ₆ ,4H ₂ O	
	-	NaC ₂ H ₃ O ₂ ,3H ₂ O	•
		$NaC_7H_5O_2$	• •
		NaHCO ₃ o	
		$Na_2B_4O_7$, $10H_2O$	
		Na ₂ CO ₃	
		Na ₂ CO ₃ 10H ₂ O	
		NaOH	.0040 ''
		NaC ₇ H ₈ O ₈ o	
No. 241.	DECINORMAL SODIUM HYDROXIDE		
		$H,C_2H_3O_2$	
		H ₃ BO ₃	
		$H_3C_6H_5O_7,H_2O$.0070 "
	" hydrobromic	HBBr o	.0081 "
	" hydrochloric	HCl o	.00365 "
	" hydriodic	HI o	.0128 "
		HC ₃ H ₅ O ₃	
	" malic	C ₄ H ₆ O ₅	-0067 "
	" nitric	HNO ₃ o	.осбз "
	" oxalic	$H_2C_2O_4$, $_2H_2O_2$.0063 "
	phosphoric	o the phenolphthalein	.0049 "
Ì	** phosphoric	H ₃ PO ₄ to form KH ₂ PO ₄ with methyl orange	.0098 "
	" sulphuric	H ₂ SO ₄	.0049 "
		H ₂ C ₄ H ₄ O ₆	
	_	KHC,H,O,	
		Na ₂ B ₄ O ₇ , 10H ₂ O	

^{*} To be ignited.

No. 142.	DECINORHAL IODINE SOLUTION.	One cc. is equivalent to	
•		As ₂ O ₃ 0.00495	gram.
		K ₂ SO ₂ 2H ₂ O 0.0097	٠,
		NaHSO ₃ 0.0052	"
		Na ₂ SO ₂ 7H ₂ O 0.0126	"
		Na ₂ S ₂ O ₂ 5H ₂ O 0.0248	"
		SO ₂ 0.0032	"
		H ₂ SO ₃ 0.0041	"
No. 245.	DECINORMAL SODIUM THIOSULPH	ATE SOLUTION. One cc. is equivalent to	
		Br 0.0080	gram.
		Cl	"
	Iodine	I 0.01266	"
	Iron (in ferric salts)	Fe	"
No. 230.	DECINORMAL SILVER NITRATE SO		
-		NH ₄ Br 0.0098	gram.
		NH,Cl 0.00535	
		Cl	
	Cyanogen	(CN) ₂ 0.0052	"
		HCN with indicator 0.0027	"
		HCN { to formation of precip- } 0.0054	"
	Hydrobromic acid	HBr	"
	Potassium bromide	KBr o.o119	"
	" chloride	KCl	"
	" cyanide	KCN with indicator 0.0065	"
		KCN { to formation of precip- } o.or30	"
	Sodium bromide	NaBr	"
	" chloride	NaCl 0.00585	"
No. 201.	DECINORMAL POTASSIUM BICHROS	MATE SOLUTION.† One cc. is equivalent t	0
	Ferrous carbonate	FeCO ₂ 0.0116	zram.
		Fe ₂ O ₃ o.0080	".
	Ferrous oxide	FeO	"
	" sulphate	FeSO, 0.0152	"
		FeSO ₄ ,7H ₂ O 0.0278	"
		Fe 0.0056	"
No. 220.	DECINORMAL POTASSIUM PERMAN	GANATE SOLUTION. One cc. is equivalen	t to
	Oxalic acid	$H_2C_2O_4$, $2H_2O_2$ 0.0063	gram,
	and to same weights for iron	salts as given under N/10 K ₂ Cr ₂ O ₇ .	

Use potassium chromate solution as an indicator, or add till precipitate appears.
 † Use a freshly prepared solution of potassium ferricyanide as an indicator, applying a drop of titrated solution to a drop of indicator on a white surface.

The following table from Talbot * shows the reactions of the common indicators used in acidimetry:

Indicator.	Reaction with Acids.	Reaction with Alkalies.	Use with Carbonic Acid in Cold Solution.	Use with Carbonic Acid in Hot Solution.	Use with Ammonium Salts.	Use with Organic Acid.
Litmus. Methyl orange. Phenolphthalein. Lacmoid. Cochineal Rosolic acid Alizarine.	Pink Colorless Purple-red Purple-red Yellow	Blue Yellow Pink Blue Blue Pink Red	Unreliable Reliable Unreliable Unreliable Reliable Unreliable Unreliable	Reliable Unreliable Reliable Reliable Reliable Reliable Reliable	Reliable Reliable Unreliable Reliable Reliable Unreliable Reliable	Reliable Unreliable Reliable Unreliable (?) Unreliable Unreliable† Reliable

^{*} Talbot, Quantitative Analysis, page 75.

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[†] Reliable with oxalic acid.

CHAPTER III.

FOOD, ITS FUNCTIONS, PROXIMATE COMPONENTS, AND NUTRITIVE VALUE.

Nature and General Composition.—Food is that which, when eaten, serves by digestion and absorption to support the functions and powers of the body, by building up the material necessary for its growth and by supplying its wastes. The raw materials that constitute our foodsupply are not all available for nourishment, but often contain a proportion of inedible or refuse matter, which it is customary to remove before eating, such as the bones of fish and meat, the shells of clams and oysters. eggshells, the bran of cereals, and the skins, stones, and seeds of fruits and vegetables. The proximate components which make up the edible portion of food include in general water, fat, various nitrogenous bodies consisting chiefly of proteins, carbohydrates, organic acids, and mineral matter. Of these water is hardly to be considered as a nutrient, though it plays an important part in nearly all foods as a diluent and solvent. The fats, proteins, and carbohydrates all contribute in varying degree to the supply of fuel for the production of heat and energy. Besides this universal function, the fats and the carbohydrates serve especially to furnish fatty tissue in the body, while the proteins are the chief source of muscular tissue.

Liebig's classification of foods into nitrogenous, or flesh formers, and non-nitrogeneous, or heat generators, is now no longer accepted as strictly logical, in view of the well-known fact that the nitrogenous materials, besides building up the body, aid in supplying the wastes and yielding energy, and may even be converted into fats or carbohydrates, while the non-nitrogenous aid in furnishing tissue growth in addition to serving as fuel.

THE FAT OF FOOD.—Fats are the glycerides of the fatty acids, the characteristics of the various edible fats and oils being treated of under

their appropriate headings elsewhere. Fat in human food is supplied by milk and its products, by the adipose tissue of meat, and in slight extent by the oil of cereals and by the edible table oils. The term "ether extract" is sometimes used in stating the results of the analysis of foods and this includes other substances than fat which when present are extracted by ether, such as chlorophyl and other coloring matters, lecithin, alkaloids, etc.

NITROGENOUS COMPOUNDS AND THEIR CLASSIFICATION.—These substances may for convenience be grouped as follows:

A Proteins, B Amino-acids and Amides, C Alkaloids, D Nitrates, E Ammonia, and F Lecithin.

A. PROTEINS.—This term includes a large number of nitrogenous bodies consisting, according to our present knowledge, essentially of combinations of α -amino-acids and their derivatives. Proteins in one form or another exist in nearly all natural foods both animal and vegetable. The terms "proteids" or "albuminoids" were formerly used generically as synonymous with "protein" to include all nitrogenous bodies of this group, but recently a joint committee on protein nomenclature of the American Physiological Society and the American Society of Biological Chemists recommended that the word "proteid" be abandoned; that "protein" be used to designate the entire group; and that the word "albuminoids" be restricted to a subgroup of proteins. A committee of the Physiological Society of England also made the same recommendation as to the use of the term protein. The classification and most of the definitions here given are those adopted by the American committee.*

Proteins available for food are supplied chiefly by the flesh of meat and fish, by milk, cheese, and eggs, and in the vegetable kingdom by seeds, nuts, and vegetables, especially the legumes. The proportion of crude protein, often designated merely as "protein," is commonly estimated by multiplying by 6.25 the percentage of nitrogen found in the material analyzed. This is done on the assumption that all of the nitrogen present in the substance belongs to protein containing 16 per cent of nitrogen.

There is no marked distinction in chemical constitution between animal and vegetable proteins, although some of the types have as yet been found only in one or the other kingdom. All proteins are insoluble in pure alcohol or in ether. A few are soluble in water but most are not. Nearly all are soluble in very dilute acids or alkalies, while all are decomposed by boiling with concentrated mineral acids or concentrated caustic alkalies. All proteins are lævo-rotary with polarized light.

^{*} Am. Jour. Phys., 21, 1908, p. xxvii.

Qualitative Test for Proteins.—Xanthoproteic Reaction.—Concentrated nitric acid added to a solution of a protein may or may not form a precipitate. It, however, produces a yellow coloration on boiling. Addition of ammonia in excess turns the precipitate or liquid yellow or orange.

Millon's Reaction.—Millon's reagent No. 184, page 30, when added to a protein solution produces a white precipitate, which becomes brick-red on heating. Sodium chloride prevents the reaction. Various organic substances are precipitated by Millon's reagent, but these precipitates do not turn red on heating.

Biuret Reaction.—If a solution of a protein in dilute sulphuric acid be made alkaline with potassium or sodium hydroxide and very dilute copper sulphate be added, a reddish to violet coloration is produced, similar to that formed if biuret* be treated in the same way, hence the name. An excess of copper sulphate should be avoided lest its color obscure that of the reaction.

In solutions which are strongly colored this reaction is of little use unless modified as follows: A considerable quantity of the dilute copper sulphate solution is added to the solution made alkaline with a large excess of potassium hydroxide, and then solid potassium hydroxide is dissolved to almost complete saturation in the solution. The mixture is then shaken with about one half its volume of strong alcohol. On standing the alcohol separates as a clear layer of a violet or crimson color if proteins are present.

I. THE SIMPLE PROTEINS.—Protein substances which yield only α -amino acids or their derivatives on hydrolysis.

Although no means are at present available whereby the chemical individuality of any protein can be established, a number of simple proteins have been isolated from animal and vegetable tissues which have been so well characterized by constancy of ultimate composition and uniformity of physical properties that they may be treated as chemical individuals until further knowledge makes it possible to characterize them more definitely.

(a) Albumins.—Simple proteins soluble in pure water and coagulable by heat.

Examples.—Seralbumin of blood and other animal fluids; lactalbumin of milk; leucosin of the seeds of wheat, rye, and barley; legumelin of leguminous seeds.

^{*} Biuret is the substance formed by heating urea to 160° according to the following reaction:

Coagulation.—Animal albumins usually coagulate at about 75°; vegetable albumins at about 65°.

Miscellaneous Reactions.—Very dilute acids precipitate albumins with the aid of heat. Nitrate of mercury (in dilute nitric acid) precipitates albumins from their solutions; also Mayer's solution acidified with acetic acid. They are precipitated by saturation with ammonium sulphate.

These reactions are not, however, characteristic of the group.

(b) Globulins.—Simple proteins insoluble in pure water, but soluble in neutral solutions of salts of strong bases with strong acids.

Examples.—Myosin of muscle substance; legumin of leguminous seeds; amandin of almonds.

Qualitative Tests.—Globulins are precipitated from their solution by dialysis or dilution. Albumins are not thus precipitated.

(c) Glutelins.—Simple proteins insoluble in all neutral solvents, but readily soluble in very dilute acids and alkalies.

Examples.—Glutenin of wheat is the only well defined protein of this group.

(d) Prolamins.—Simple proteins soluble in relatively strong alcohol (70-80 per cent), but insoluble in water, absolute alcohol, and other neutral solvents.

Examples.—Gliadin of wheat; zein of maize; hordein of barley. Found as yet only in the seeds of cereals.

The use of appropriate prefixes will suffice to indicate the origin of compounds of sub-classes a, b, c, and d, as for example: ovoglobulin, myalbumin, etc.

(e) Albuminoids.—Simple proteins which possess essentially the same chemical structure as the other proteins, but are characterized by great insolubility in all neutral solvents.

Examples.—Keratins of hair, nails, hoofs, horn, feathers, etc.; elastin of connective tissues; collagen of connective tissues and cartilage; fibroin and sericin of raw silk. No albuminoids have yet been discovered in plants.

Gelatin is usually regarded as an albuminoid but does not come strictly within the requirements of the above definition. It is an artificial derivative of collagen and is formed from it by boiling with water or subjecting to steam under pressure. It is prepared from bones and other animal parts, and is insoluble in cold, but soluble in hot water. When the hot water solution containing one per cent or more of gelatin cools, it forms a jelly. By prolonged boiling the gelatinizing power is lost. Aqueous solutions are strongly lævo-rotary.

Gelatin in common with most proteins is precipitated from its solution by mercuric chloride, picric acid, and tannic acid. It is readily distinguished from soluble proteins, in that it is not precipitated from its solution by lead acetate, nor by most of the metallic salts that throw down proteins.

(f) Histones.—Soluble in water and insoluble in very dilute ammonia, and, in the absence of ammonium salts, insoluble even in an excess of ammonia; yield precipitates with solutions of other porteins, and a coagulum on heating, which is easily soluble in very dilute acids. On hydrolysis they yield a large number of amino-acids, among which the basic ones predominate.

Examples.—Thymus histone. Not found in plants.

(g) Protamins.—Simpler polypeptides than the proteins included in the preceding groups. They are soluble in water, uncoagulable by heat, have the property of precipitating aqueous solutions of other proteins, possess strong basic properties, and form stable salts with strong mineral acids. They yield comparatively few amino-acids, among which the basic amino-acids greatly predominate.

Examples.—Salmin, clupein, and other protamins of fish spermatozoa. Not found in plants.

- II. CONJUGATED PROTEINS.—Substances which contain the protein molecule united to some other molecule or molecules otherwise than as a salt.
- (a) Nucleoproteins.—Compounds of one or more protein molecules with nucleic acid.

Examples.—The nucleins formed by pepsin digestion.

(b) Glycoproteins.—Compounds of the protein molecule with a substance or substances containing a carbohydrate group other than a nucleic acid.

Examples. - Mucins; ovomucoid; ovalbumin.

(c) Phosphoproteins.—Compounds of the protein molecule with some yet undefined phosphorus-containing substance other than a nucleic acid or lecithins.

Examples.—Casein of milk; vitellin of egg yolk.

(d) Haemoglobins.—Compounds of the protein molecule with haematin or some similar substance.

Example.—Oxyhaemoglobin of red blood corpuscles.

(e) Lecithoproteins.—Compounds of the protein molecule with lecithins, (lecithans, phosphatides).

Examples.—Lecithalbumin; lecithin-nucleovitellin.

III. DERIVED PROTEINS.

- 1. PRIMARY PROTEIN DERIVATIVES.—Derivatives of the protein molecule, apparently formed through hydrolytic changes which involve only slight alterations of the molecule.
- (a) Proteans.—Insoluble products which apparently result from the incipient action of water, very dilute acids or enzymes.

Examples.—Edestan; blood fibrin; insoluble myosin.

(b) Metaproteins.—Products of the further action of acids or alkalies, whereby the molecule is so far altered as to form products soluble in very weak acids and alkalies, but insoluble in neutral fluids.

Examples.—Acid albumin; alkali albumin.

This group will thus include the familiar "acid proteins" and "alkali proteins," not the salts of proteins with acids.

(c) Coagulated Proteins.—Insoluble products which result from (1) the action of heat on their solutions, or (2) the action of alcohol on the protein.

Examples.—Albumin coagulated by heat or alcohol.

- 2. SECONDARY PROTEIN DERIVATIVES. Products of the further hydrolytic cleavage of the protein molecule.
- (a) Proteoses.—Soluble in water, uncoagulated by heat, and precipitated by saturating their solutions with ammonium or zinc sulphate.

As thus defined this term does not strictly cover all the protein derivatives commonly called proteoses, e.g. heteroproteose and dysproteose.

Subdivision of the Proteoses.—According to the proteins from which they are derived the proteoses may be designated albumose, from albumin, globulose, from globulin, vitellose, from vitellin, caseose, from casein, etc.

Proteoses are subdivided into proto-proteose, soluble in water (both cold and hot) or in dilute salt solutions, but precipitated by saturation with salt; hetero-proteose, insoluble in water, and deutero-proteose, soluble in water, but not precipitated by saturation with salt.

Vegetable proteoses are sometimes called phyt-albumoses.

Qualitative Tests.—Besides responding to the biuret reaction (p. 41) proteoses are precipitated by nitric acid, the precipitate being soluble on heating, but reappearing on cooling.

Proto-proteose is precipitated from its solution by mercuric chloride and copper sulphate; hetero-proteose is precipitated by mercuric chloride, but not by copper sulphate.

(b) Peptones.—Soluble in water, uncoagulated by heat, and not precipitated by saturating their solutions with ammonium sulphate.

Qualitative Tests.—Besides giving the biuret reaction, peptones are precipitated from their solution by tannic acid, picric acid, phosphomolybdic acid, and by sodium phosphotungstate acidified by acetic, phosphoric, or sulphuric acid.

Peptones are the only soluble proteins not precipitated by saturation with ammonium sulphate. The following table, showing the reaction of proteoses and peptones, is due to Halliburton:*

Variety of Protein.	Hot and Cold Water.	Hot and Cold Saline Solutions, e.g., 10% NaCl.	Satura- tion with NaCl or MgSO ₄ .	Satura- tion with (NH ₄) ₂ SO ₄	Nitric Acid.	Copper Sulphate.	Copper Sulphate and Caustic Potash.
Proto- albumose	Soluble	Soluble	Precipi- tated	Precipi- tated	Precipitated in cold; pre- cipitate dis- solves with heat and re- appears on cooling	Precipi- tated	Rose-red color (biu- ret reac- tion)
Hetero- albumose	Insoluble; i.e. precipitated by dialysis from saline solutions	Soluble; part- ly precipita- ted, but not coagulated on heating to 65° C.	Precipi- tated	Precipi- tated	Ditto	Precipi- tated	Ditto
Deutero- albumose	Soluble	Soluble	Not pre- cipitated	Precipi- tated	This reaction occurs only in presence of excess of salt	Not pre- cipitated	Ditto
Peptone	Solub ie	Soluble	Not pre- cipitated	Not pre- cipitated	Not pre- cipitated	Not pre- cipitated	Ditto

(c) Peptides.—Definitely characterized combinations of two or more amino-acids, the carboxyl group of one being united with the amino group of the other, with the elimination of a molecule of water.

The peptones are undoubtedly peptides or mixtures of peptides, the latter term being at present used to designate those of definite structure.

- B. AMINO-ACIDS, AMIDES, AND ALLIED PRODUCTS.—Under this head are included products derived from acids or bases, the radicles of which replace one or more of the hydrogen atoms in ammonia. The most common bodies of this class occurring in food products are:
- (1) Cholin (C₅H₁₅NO₂), found in the muscular tissue of cattle and in yolk of eggs, also in certain fungi.
- (2) Betaine (C₅H₁₁NO₂), found in certain mollusks, as, for instance, the mussel, in putrefying fish, and (in the vegetable kingdom) in beets and hops. It is formed by the oxidation of cholin.
- (3) Asparagin (C₄H₈N₂O₃), found in the shoots of asparagus, lettuce, peas and beans, and in the root of the marshmallow. It may be crystal-

^{*} Chemical Physiology and Pathology, page 131.

lized out from the expressed juice of the asparagus shoots by evaporation, after having removed the albumin by coagulation (by boiling) and by filtration.*

Asparagin when heated with alkalies forms ammonia, and with acids forms ammonium salts. Freshly prepared copper hydroxide is dissolved by an aqueous solution of asparagin by the aid of heat. If sections of vegetable tissues containing asparagin are placed in alcohol, crystals of asparagin are formed in such a manner as to be detected under the microscope.†

Closely allied to the amides are the flesh bases of meat, chief among which are creatin ($C_4H_9N_3O_2$), creatinin ($C_4H_7N_3O$), derived from creatin by the action of mineral acids and existing in some fish, carnin ($C_7H_9N_4O_3$), and xanthin ($C_5H_4N_4O_2$).

- C. ALKALOIDAL NITROGEN.—Alkaloids do not naturally occur in foods, with the exception of tea, coffee, and kola-nuts, which contain caffeine, and cocoa, which contains theobromine.
- D. NITROGEN AS NITRATES.—Foods in their natural condition rarely contain nitrates. Meats cured with saltpetre furnish the most common instance of nitrates in food. Nitrates are tested for by extracting the sample with water, and treating the extract with ferrous sulphate and sulphuric acid in the usual manner.
- E. NITROGEN AS AMMONIA. Ammonia occurs very sparingly in food, unless the latter has undergone some form of decomposition. In ripened cheese and in sour milk one sometimes finds it in minute quantities. Its presence is tested for by distilling the finely divided sample in water free from ammonia, and testing the distillate with Nessler's reagent.
- F. LECITHIN.—This substance (C₄₄H₉₀NPO₉) is a phosphorized fat, and forms a part of the cell material in certain animal and vegetable foods. It is found in considerable quantity in the yolk of egg, and, in traces, in cereals, peas, and beans. It is a yellowish-white solid, soluble in ether and alcohol. Treated with water it swells up, forming an opalescent solution or emulsion, from which it is precipitated by salts of the alkali metals.

THE CARBOHYDRATES AND THEIR CLASSIFICATION.—The carbohydrates supplied by food are milk sugar and the various sugars, starches, and gums from plant juices, cereals, fruits, and vegetables. Carbohydrates are generally understood as being compounds of carbon, hydrogen, and oxygen, the last two elements being present in the proportion in

^{*} Zeits. für analytische Chemie, 22, page 325.

[†] Wiley, Principles of Agric'l Analysis, Vol. III. p. 427.

which they occur in water. They are divided into three main classes, as follows:

- A. The Glucose Group, or Monosaccharids (C₆H₁₂O₆), including dextrose, levulose, and galactose.
- B. The Cane Sugar Group, or Disaccharids (C₁₂H₂₂O₁₁), including cane sugar, milk sugar, and maltose.
- C. The Cellulose Group (C₆H₁₀O₅), including starch, cellulose, dextrin, gums, etc.

Closely allied to the carbohydrates, if not actually belonging to them, are *inosite* (C₆H₁₂O₆), occurring in muscular tissue, and *pectose*, found in green fruits and vegetables.

THE ORGANIC ACIDS.—These acids are minor though important constituents of foods. From their conversion into carbonates within the body, they are useful in furnishing the proper degree of alkalinity to the blood and to the various other fluids, besides being of particular value as appetizers. They exist in foods both in the free state and as salts. Acetic acid is supplied by vinegar; lactic acid by milk, fresh meat, and beer; citric, malic, and tartaric acids by the fruits.

MINERAL OR INORGANIC MATERIALS.—These substances occur in food in the form of chlorides, phosphates, and sulphates of sodium, potassium, calcium, magnesium, and iron, and are furnished by common salt, as well as by nearly all animal and vegetable foods. The inorganic salts are necessary to supply material for the teeth and bones, besides having an important place in the blood and in the cellular structure of the entire body.

FUEL VALUE OF FOOD.—In order to express the capacity of foods for yielding heat or energy to the body, the term fuel value is commonly used. By the fuel value of a food material is meant the amount of heat expressed in calories equivalent to the energy which we assume the body could obtain from a given weight of that food material, if all of its nutritents were thoroughly digested, a calorie being the amount of heat required to raise a kilogram of water 1° C. This definition applies to what is known as the large calorie, which is one thousand times as large as the small calorie. Large calories are meant wherever the term occurs in this volume. The fuel value, or, as it is sometimes called, "heat of combustion," may be determined experimentally with a calorimeter, or may be calculated by means of factors based on the result of many experiments showing the mean values for protein, fats, and carbohydrates.

The Bomb Calorimeter.*—This apparatus in its most approved form,

^{*}U. S. Dept. of Agric., Off. of Exp. Sta., Bul. 21, pp. 120-126.

Fig. 13, consists of a water-tight, cylindrical, platinum lined, steel bomb, adapted to hold in a capsule the substance whose heat is to be determined, and containing also oxygen under pressure. This bomb is immersed in water contained in a metal cylinder, which is in turn placed inside of concentric cylinders containing alternately air and water. The heat for igniting the substance is supplied by the electric current passing through wires to the interior of the bomb and acting upon a cleverly devised mechanism therein. The heat developed by the ignition is measured by

Fig. 13.—Bomb Calorimeter of Hempel and Atwater

the rise in temperature of the water surrounding the bomb, as indicated by a very delicate thermometer graduated to hundredths of a degree, certain corrections being made, as, for instance, for the heat absorbed by the metal of the apparatus. A mechanical stirrer serves to equalize the temperature of the water surrounding the bomb.

Calculation of Fuel Value.—By reason of its great expense the calorimeter is beyond the reach of many laboratories, and on this account the expression of fuel values by calculation is the most common method employed. For this the factors of Rubner are generally used, in accordance with which the amount of energy in one gram of each of the three principal classes of nutrients are, for carbohydrates 4.1, for protein 4.1, and for fats 9.3. Expressed in pounds, each pound of carbohydrate or protein has a fuel value of 1860 calories, while each pound of fat has a fuel value of 4220 calories.

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- Bul. 63. Description of a New Respiration Calorimeter and Experiments on the Conservation of Energy in the Human Body. By W. O. Atwater and E. B. Rosa. Pp. 94.
- Bul. 68. A Description of Some Chinese Vegetable Food Materials and their Nutritive and Economic Value. By W. C. Blasdale. Pp. 48.

- Bul. 69. Experiments on the Metabolism of Matter and Energy in the Human Body. By W. O. Atwater and F. G. Benedict, with the cooperation of A. W. Smith and A. P. Bryant. Pp. 112.
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- Bul. 85. A Report of Investigations on the Digestibility and Nutritive Value of Bread. By C. D. Woods and L. H. Merrill. Pp. 51.
- Bul. 89. Experiments on the Effect of Muscular Work upon the Digestibility of Food and the Metabolism of Nitrogen. Conducted at the University of Tennessee, 1807-1809. By C. E. Wait. Pp. 77.
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- Bul. 121. Experiments on the Metabolism of Nitrogen, Sulphur, and Phosphorus in the Human Organism. By H. C. Sherman. Pp. 47.
- Bul. 126. Studies on the Digestibility and Nutritive Value of Bread at the University of Minnesota in 1900–1902. By Harry Snyder. Pp. 52.
- Bul. 129. Dietary Studies in Boston and Springfield, Mass., Philadelphia, Pa., and Chicago, Ill. By Lydia Southard, Ellen H. Richards, Susannah Usher, Bertha M. Terrill, and Amelia Shapleigh. Pp. 103.
- Bul. 132. Further Investigations Among Fruitarians at the California Agricultural Experiment Station. By M. E. Jaffa. Pp. 81.
- Bul. 136. Experiments on the Metabolism of Matter and Energy in the Human Body, 1900-1902. By W. O. Atwater and F. G. Benedict. Pp. 357.
- Bul. 143. Studies on the Digestibility and Nutritive Value of Bread at the Maine Agricultural Experiment Station, 1899–1903. By C. D. Woods and L. H. Merrill. Pp. 77.

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- Bul. 150. Dietary Studies at the Government Hospital for the Insane, Washington, D. C. By H. A. Pratt and R. D. Milner. Pp. 170.
- Bul. 152. Dietary Studies with Harvard University Students. By E. Mallinckrodt, jr. Pp. 63.
- Bul. 156. Studies on the Digestibility and Nutritive Value of Bread and Macaroni at the University of Minnesota, 1903–1905. By Harry Snyder. Pp. 80.
- Bul. 159. A Digest of Japanese Investigations on the Nutrition of Man. By K. Oshima. Pp. 224.
- Bul. 162. Studies on the Influence of Cooking upon the Nutritive Value of Meats at the University of Illinois, 1903–1904. By H. S. Grindley and A. D. Emmett. Pp. 230.
- Bul. 175. Experiments on the Metabolism of Matter and Energy in the Human Body. 1903–1904. By F. G. Benedict and R. D. Milner. Pp. 335.
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- Bul. 187. Studies of the Digestibility and Nutritive Value of Legumes at the University of Tennessee, 1901–1905. By C. E. Wait. Pp. 55.
- Bul. 193. Studies of the Effect of Different Methods of Cooking upon the Thoroughness and Ease of Digestion of Meat at the University of Illinois. By H. S. Grindley. Pp. 100.
- Bul. 208. The Influence of Muscular and Mental Work upon Metabolism and the Efficiency of the Human Body as a Machine. By F. G. Benedict.

CHAPTER IV.

GENERAL ANALYTICAL METHODS.

Extent of a Proximate Chemical Analysis.—For purposes of studying the proximate composition of food for its dietetic value, it is nearly always necessary to make determinations of moisture, ash, fat, total nitrogen, and carbohydrates (when present), as well as of the fuel value. In some cases it may be desirable to proceed further, to make an analysis of the ash, for instance, to separate, at least into classes, the various nitrogenous bodies, especially in flesh foods, and perhaps to subdivide the starch, sugar, gums, and cellulose or crude fiber that make up the carbohydrates in the case of cereals.

An analysis is considered complete whenever the purpose for which the examination has been made has been accomplished, and on that purpose depends solely the extent to which the various compounds present shall be subdivided and determined. Such a subdivision may be extended almost indefinitely. For example, a milk analysis may consist simply in the determination of the total solids and (by difference) the water. Again, it may be desirable to divide the milk solids into fat and solids not fat, and in some cases to carry the subdivision still farther and separate the solids not fat into casein, albumin, milk sugar, and ash.

Determinations of one or more of the proximate components natural to food are frequently of great service in proving their purity or freedom from adulteration. For the latter purpose, especially in such foods as milk, vinegar, oils, and fats, the determination of specific gravity is also an important factor. Special methods of a peculiar nature are often necessary in the examination of particular foods, and such methods will be treated subsequently under the appropriate headings. In the present chapter only such general methods as are applicable to a large variety of cases will be discussed.

Expression of Results of a Proximate Analysis.—However complete the division of the various proximate compounds or classes of compounds

which the analyst sees fit to make, the results of his various determinations in a proximate analysis are expected to aggregate about 100%. If every determination be directly made, the result will rarely be exactly 100, but the precision of the work is apt to be judged by its approach to 100.

It is often the custom to determine certain compounds or classes of compounds by difference. Thus in cereals moisture, proteins, fat, crude fiber and ash may be determined by the regular analytical methods, and by subtracting their sum from 100 the difference may be expressed as "nitrogen-free extract" or carbohydrates. It has long been customary in food analysis to calculate the protein by multiplying the total nitrogen by the factor 6.25, and on this basis analyses of thousands of animal and vegetable foods have been made. While the figure thus obtained is an arbitrary one, being at best but a rough approximation of the amount of protein present, yet for many reasons there is much to commend this practice of reporting results. In the first place, in most cases it actually does approach the truth. Again, the nitrogenous ingredients of many foods are so numerous and varied, that for the every-day study of dietaries and food values it would be well-nigh impossible with our present knowledge to subdivide these compounds with any degree of accuracy, and especially with uniformity between different chemists, to say nothing of the time involved.

From the fact that so many valuable analyses have already been expressed on the basis of $N \times 6.25$ for protein, the advantage of comparison with the results thus recorded would seem to be in itself a good reason for continuing the practice, especially until a factor that gives better average results can be adopted. By recording the actual nitrogen found as well as the "protein," old results may at any time be recalculated under new conditions, if found desirable.

In flesh foods, when carbohydrates are known to be absent, the total protein may conveniently be determined by difference. Rather more progress has been made in the separation of the nitrogenous compounds of meats than of the vegetables and cereals, though the methods are by no means accurate or uniform.

Most of the recorded analyses of vegetable foods express the carbohydrates as a whole without attempting to subdivide them, at least further than possibly to express the crude fiber or cellulose separately. A much more intelligible idea of the dietetic value of these foods would be gained by a further separation into starch and sugars.

Preparation of the Sample.—It is at the outset of the utmost importance in all cases that a strictly representative portion of the food to be examined should be submitted to analysis. All refuse matter, such as bones, shells, bran, skins, etc., are removed as completely as possible from the edible portion and discarded.

If the composition of the entire mass cannot be made homogeneous throughout, it may be best to select from various portions in making up the sample for analysis, in order to represent as fair an average of the whole as possible.

Finally the sample, if solid or semi-solid, should be divided as finely as possible, by chopping, shredding, pulping, grinding, or pulverizing according to its nature and consistency.

For disintegrating such substances as vegetables and meats for analysis, the common household rotary chopping-machine is admirably adapted. For pulverizing cereals, tea, coffee, whole spices, and the like, the mortar and pestle may be used, or a rotary disk mill or spice-grinder.

Specific Gravity or Density of Liquids.—Where formerly it was customary to compare the density of liquids with that of water at 4° C. (its maximum density) it is now more common to refer to water at 15.5° C. or 20° C., making the determination at that temperature. A common form of expressing the temperature of the determination and the temperature of the standard volume of water with which that of the substance is to be compared, is the employment of a fraction, the numerator of which expresses the temperature of the determination and the denominator

that of the standard volume of water, as $\frac{15.5^{\circ}}{4^{\circ}}$, $\frac{15.5^{\circ}}{15.5^{\circ}}$, $\frac{100^{\circ}}{15.5^{\circ}}$, $\frac{4^{\circ}}{4^{\circ}}$ C.* When extreme accuracy in the determination of density is required, the pycnometer or Sprengel tube should be employed.

The Hydrometer.—This instrument furnishes the most convenient and ready means of determining the density of liquids where extreme nicety is not required. If well made and carefully adjusted, the hydrometer may be depended on to three decimal places, but before relying on its accuracy, it should be tested by comparison with a standard instrument, or with the pycnometer.

The liquid whose density is to be determined is contained in a jar whose inner diameter should be at least $\frac{3}{3}$ larger than that of the spindle-

^{*} Unless otherwise stated, all specific gravities in this volume are assumed to be expressed on the basis of $\frac{15.5^{\circ}}{15.5^{\circ}}$

bulb, and the temperature of the liquid should be exactly 15.5° when the reading is taken.

For best results for use with liquids of varying densities, the laboratory should be furnished with a set of finely graduated hydrometers, each limited to a restricted part of the scale, together with a universal hydrometer coarsely graduated, covering the entire range, to show by preliminary test which of the special instruments should be used.

A convenient set of seven such hydrometers are graduated as follows: 0.700-0.850, 0.850-1.000, 1.000-1.200, 1.200-1.400, 1.400-1.600, 1.600-1.800, 1.800-2.000, while the universal hydrometer has a scale extending from 0.700 to 2.000. Another less delicate set of three only has one for liquids lighter than water and two for heavier liquids. Some instruments have thermometers in the stem. Others require a separate thermometer.

The Westphal Balance (Fig. 14).—This instrument consists of a scale-beam fulcrumed upon a bracket, which in turn is upheld by a supporting pillar. The scale-beam is graduated into ten equal divisions. From a hook on the outer end of the beam hangs a glass plummet provided with a delicate thermometer, the beam being so adjusted that when the dry plummet hangs in the air, the beam is in exact equilibrium, i.e., perfectly horizontal as shown by the indicator on its inner end. If the large rider be placed on the same hook as the plummet and the latter immersed in distilled water of the standard temperature at which the determinations are to be made (say 15.5° C.), the scale-beam should again be in equilibrium if the instrument is accurately adjusted. As commonly made, the weight of the plummet including the platinum wire to which it is attached amounts to 15 grams, and the displacement of its volume to 5 grams of distilled water at 15.5° C., the normal temperature on which the determinations are based. Thus the unit (or largest) rider should weigh 5 grams, while the others weigh 0.5, 0.05, and 0.005 gram respectively.

If, instead of distilled water, the plummet be immersed in the liquid whose density is to be determined, the position of the riders on the scale-beam, when so placed as to bring the same into equilibrium, and when read in the order of their relative size (beginning at the largest), indicates directly the specific gravity to the fourth decimal place.

If the liquid is lighter than water, the large rider is first placed in the notch where it comes closest to restoring the equilibrium of the beam, but with the plummet still underbalanced. The rider next in size is then applied in a similar manner, and, unless equilibrium is exactly re-

stored, the third and the fourth riders successively. If it happens that two riders should occupy the same position on the beam, the smaller is suspended from the larger.

If the liquid is heavier than water, the method employed is the same except that one of the largest or unit riders is in this case always hung from the hook which supports the plummet, while the others cross the

Fig. 14.—The Westphal Balance.

beam at the proper points. If carefully made and adjusted, the Westphal balance is capable of considerable accuracy.

A delicate analytical balance can be used in place of the less carefully adjusted Westphal instrument, by hanging the Westphal plummet from one of the scale-hooks of the same, and employing a fixed support for the glass jar that holds the liquid in which the plummet is to be immersed. The support is so arranged that the scale-pan below it can move freely without coming in contact with it. This arrangement is shown in Fig. 15.

The Pycnometer, or Specific-gravity Bottle. - Fig. 16 shows the two

forms of pycnometer commonly made. The plain form has a ground-glass stopper with a capillary passage through it, the other has a fine thermometer connected with the stopper and a capillary side tube provided with a ground hollow cap. Both are made in different sizes to hold respectively 10, 25, 50, and 100 grams of distilled water at the standard

Fig. 15.—The Analytical Balance Arranged for Determining Specific Gravity with the Westphal Plummet,

temperature. It is convenient to have a counterweight for each pycnometer as fitted with its stopper, thus avoiding much trouble in calculation. The calculation of results is simplified also if the pycnometers are accurately constructed to contain exactly the weight of distilled water which they purport to contain at the standard temperature, but it is rather difficult to procure such instruments, especially of the form furnished with the thermometer. Most instruments hold approximately the amount specified, the exact net weight of distilled water which they hold at standard temperature being found by careful test and kept on record. In determining the density of a liquid, the pycnometer is carefully filled with it at a temperature below the standard, the stopper carefully inserted, and the bottle wiped dry. Care should be taken that the liquid completely fills the bottle and is free from air-bubbles. The net weight of the liquid is then taken

on the balance, when the temperature has reached the standard (say 15.5° C.), being careful to wipe off the excess of liquid that exudes from the capillary due to expansion. The net weight of the liquid is divided by that of the same volume of distilled water, previously ascertained under the same conditions at the same temperature, the result being the density of the liquid.

The pycnometer with thermometer attachment is obviously susceptible of a greater degree of accuracy than the other form, since the temperature of the liquid, even though 15.5° C. at the start, soon rises.

Fig. 16.—Types of Pycnometer.

The writer prefers to use the pycnometer provided with the thermometer, but without the hollow cap that covers the capillary side tube, unless liquids like strong acids are to be operated on, that might otherwise injure the balance. By keeping the liquid to be tested for some time in a refrigerator, it acquires a temperature of from 10 to 12° C. It is then transferred in the regular manner to the pycnometer and the thermometer-stopper inserted (but not the hollow cap) and the bottle wiped dry. There is ample time to adjust the balance-weights with extreme care while the temperature of the liquid is rising, leisurely wiping off

at intervals with a soft towel the excess that exudes from the capillary tube, the final weight of the dry bottle and contents being made at the exact temperature of 15.5° C.

In taking the tare or adjusting the counterweight of a specific-gravity bottle, the latter should be perfectly clean and dry. It had best be rinsed first with water, then with alcohol, and finally with ether, all traces of the latter being removed by a current of dry air, or otherwise, before weighing.

In making successive determinations of density of a number of different liquids with the same pycnometer, it is sufficient to rinse the bottle once with a little of the liquid to be tested before making each determination, when the various liquids are miscible. When the liquids are immiscible, the bottle should be carefully cleaned in the manner described in the previous paragraph before making each test.

The Sprengel Tube.—The Sprengel tube is a variety of pycnometer useful when only a small quantity of the liquid to be tested is available.

It is susceptible of great accuracy. It consists of a U-shaped tube (Fig. 17), each branch of which terminates in a horizontal capillary tube bent outward. One of the capillaries, b, has a mark m thereon and has an inner diameter of about 0.5 mm. The diameter of the other capillary, a, should not exceed 0.25 mm. The liquid at room temperature is aspirated into the tube so as to fill it completely, the end b being dipped in the liquid while suction is applied at the end a. The tube is then placed in a beaker of water kept at the standard temperature, the beaker being of such size that the capillary ends rest on the edge. The temperature of the liquid in the tube may be assumed to be constant

Fig. 17.—Sprengel Tube when there is no further movement due to contractor Determining Spetion in the larger capillary end, b. The meniscus of the liquid, when cooled, should not be inside the mark m, and is brought exactly to the mark by applying a piece of bibulous paper to the other end, a. If a drop or two of liquid has to be added, this may be done by applying to the end a a glass rod dipped in the liquid. When exactly adjusted, the whole is wiped dry and quickly weighed, hung from the arm of the analytical balance. To avoid evaporation by contact with the air, the ends of the capillaries are sometimes ground to receive hollow glass caps not shown in the figure.

Determination of Moisture.—The moisture is usually calculated from the weight of dry residue left after driving out all the water by evaporation from a weighed portion of the sample, using generally from 1 to 10 grams in a tared dish. Some substances readily part with their water at 100°; others, again, require a much higher temperature or an extremely long heating. In general the highest possible degree of heat should be employed that will not affect the other constituents. Certain saccharine products should theoretically be dried at a temperature not exceeding 70° on account of the dehydration of some of the sugars at higher temperatures. On the contrary, where readily decomposable organic matter is known to be absent and the character of the substance will permit, it is sometimes possible to employ temperatures considerably above 100° for quick drying.

It is not always safe to assume that water is the only substance evaporated on drying. Thus spices and other products containing essential oils give off appreciable quantities of these oils when dried at 100°.

As it is rarely possible to attain a temperature higher than 98° in the water-oven, a gas-heated air-oven of the general type of that described on page 23, with ready means for controlling the temperature, is best for general moisture determinations in the food laboratory.

Platinum dishes like those described on page 133 are admirable for nearly all moisture determinations, but thin dishes of porcelain, glass, or metal may be used. Thin liquids and air-dry, powdered substances may usually be weighed directly in the dish and dried, without the use of an absorbent.

With very moist substances containing much cellulose as well as water, it is often advantageous to weigh into the dish and allow to simmer for a long time on the water-bath, before drying to constant weight in the air-oven at higher temperature.

Viscous substances should generally be spread over finely divided asbestos fiber, or pieces of pumice stone, or quartz sand, which should be previously ignited and weighed with the dish, the object being to divide up the weighed portion as finely as possible for its better exposure to the heated air in the drying-oven.

Determination of Ash.—For determining the percentage of the ash or mineral matter, it is often convenient to use the portion previously weighed out and dried in obtaining the moisture, the dry residue after the second weighing being in such cases burnt in the original dish over a low flame. Or, if desired, a fresh portion of the original substance may

be air-dried or subjected to a preliminary drying in the water-bath and then burnt, taking care that there is no loss by sputtering or otherwise.

Platinum dishes will be found much the most convenient in all cases where they may be safely used. In general the shallow flat dishes described on page 133 are preferable. Where lead or tin compounds are present, or when sulphides, sulphates, or phosphates are to be burnt with reducing agents, platinum is sure to be attacked and porcelain dishes or crucibles should be used instead. Platinum is also attacked at ordinally temperatures by free chlorine and bromine, and, when ignited, by free sulphur, phosphorus, arsenic, and iodine, by sulphides and by sodium or potassium hydroxide, nitrate and cyanide. Platinum dishes are liable to injury also when used for the ignition of sulphates, and phosphates with reducing substances, or with metals present that are reduced in fusion, such as mercury, bismuth, tin, lead, zinc, antimony, etc.

The degree of heat employed in ashing should be the lowest possible to insure complete oxidation of the carbon, so as to avoid driving off certain volatile salts that are sometimes present and that would be lost if the heat were too high. At a bright red heat potassium and sodium chloride are slowly volatilized, and calcium carbonate is converted into oxide; furthermore alkali phosphates fuse about particles of carbon, protecting them from oxidation. To avoid overheating it is recommended not to allow the flame to impinge directly against the dish, but to carry out the burning on a piece of asbestos paper supported on a triangle. The asbestos also serves to distribute the heat and to protect the dish from the injurious action of the direct flame on long heating. In order to burn off the last traces of carbon, a second piece of asbestos paper may be placed over the top of the dish, or the incineration may be completed in a muffle furnace. Heating should be continued till the carbon is all oxidized, which is in most cases indicated by a white ash. It is, however, sometimes impossible to obtain a perfectly white ash, but the appearance of the ash usually indicates when all the carbon has been burnt off. It is sometimes necessary to stir the contents of the dish with a stiff platinum wire from time to time during the ignition, the wire being weighed in with the dish.

After ignition the dish is cooled in the desiccator before weighing.

If an examination of the ash for special ingredients is required, it is often necessary to burn a large portion of the sample. In this case it may be desirable to hasten the ignition by the careful use of ammonium nitrate as an oxidizing agent, or, in very refractory cases, as when sugar is present, it is well before igniting to saturate the sample with concen-

trated sulphuric acid, when the presence of sulphates in the ash is not objectionable.

Methods for the detection and determination of the various ash ingredients are considered on pages 301 to 305. Such cases as are peculiar to certain foods, like the metallic impurities that occur in canned, bottled, and preserved foods under certain conditions, will be considered in their appropriate place.

whatever it dissolves. The operation is at once repeated, the substance

ever it is necessary to exhaust a substance of its ether-soluble or alcohol-soluble ingredients, some form of continuous extraction apparatus is employed with advantage.

The Soxhlet Extractor. - This apparatus, or one of its modifications is most commonly employed for continuous extraction. shows the simplest form of the Soxhlet extractor, consisting essentially of a wide tube, A, provided with the side siphon a, a condenser, B, and a wide-mouthed flask, C, all connected together in the manner illustrated, either by soft, accurately fitting corks or by ground joints, or by mercury-sealed connections. Care in either case should be taken to have the joints perfectly tight, so as to avoid loss by leakage. The construction is such that the substance to be extracted, which is contained in the tube A, is subjected to successive treatment with freshly distilled portions of the solvent. The vapor from the solvent, boiling in the flask C, passes up through the side tube a' into the cold condenser B, where it is again reduced to liquid Fig. 18.—The Soxhlet Extractor and falls drop by drop upon the substance to be

and falls drop by drop upon the substance to be with Electric Heater.

extracted, which is confined in a suitable porous receptacle or perforated vessel in the tube A. The substance is thus allowed to macerate in the solvent till the level of the latter reaches the top of the siphon, when all of the solvent in the tube drains off into the flask C, carrying with it

being subjected to successive extractions with freshly distilled portions of the solvent, which leaves behind in the flask C whatever it dissolves. This operation of continuous extraction, when the conditions are right, goes on indefinitely without attention.

The weighed portion of the sample to be extracted (from 2 to 5 grams) is first deprived of its moisture by drying, if free from volatile oil, and then transferred to the bottom of the tube A. There are various methods of doing this. If the substance is a fluid or semi-fluid like milk, it may be taken up on an absorption-coil of fat-free filter-paper and dried (see page 135), the dried coil being transferred to the tube A. Or the sample may be weighed into a very thin glass shell (Hoffmeister's Schälchen) in which it is dried, after which the shell is wrapped in bibulous paper, crushed between the fingers into small bits, and the whole, in the form of a small packet, is placed in the tube A. Or, again, the material, if in the form of an air-dried powder, may be weighed in a tared platinum dish or watch-glass and transferred by a brush into a partly folded wrapper of filter-paper, the ends of which are afterwards closed in by folding to form a packet, which is first dried thoroughly in the oven and then placed in the tube A. The fat-free porous shells made by Schleicher & Schüll, in various sizes to fit the Soxhlet tubes, form convenient receptacles for the extraction of dry substances. The sample may in most cases be directly weighed into one of these shells after taking its tare, and the drying and extraction carried out at once.

Preliminary to conducting the extraction, the flask C, Fig. 18, is thoroughly cleaned and dried and then weighed, after which enough of the solvent reagent is poured into it to last through the period of the extraction, and the parts of the apparatus are connected.

The heater employed should be a water-bath, or, as shown in Fig. 18, an electric stove, which may be provided with a fractional rheostat for varying the amount of heat.

The degree of ebullition is so regulated as to allow the solvent to saturate the sample and siphon over into the flask C from six to twelve times an hour, the extraction being continued from two to six hours, or until all the ether-soluble material has been removed. Care should be taken also that the rate of boiling and the rate of condensation are so regulated that no appreciable loss of reagent occurs during the extraction. A strong smell of ether perceptible at the top of the condenser indicates a loss. The solvent is recovered at the end of the extraction by disconnecting the weighing flask at a time when nearly all of the solvent is in the part A and before

it is ready to siphon over. The weighing-flask is then freed from all traces of the solvent by drying first on the water-bath and then in the oven, after which it is cooled in the desiccator and weighed, the difference between this and the first weighing representing the weight of the fat or ether extract.

The Johnson Ext. actor.—This form of apparatus (Figs. 19 and 20) has the advantage of the Soxhlet extractor in that it is simpler and employs a

much smaller amount of ether. The substance is contained in the inner tube of the extractor (Fig. 19), which is closed at the lower end by one thickness each of filter paper and cheese cloth, held tightly in place by means of a linen thread wrapped several times about the tube in the constriction and tied in a fast knot. This innner tube properly prepared can be used over and over for extractions. The outer tube, also shown in Fig. 19, is of such a size that the inner tube fits loosely within it. A slight bulge on one side prevents trapping by means of the condensed solvent. The extraction flask is preferably of only 30 to 35 cc. capacity. It is attached to the extractor, as is also the extractor to the condenser tube, by means of a carefully bored cork stopper. For ordinary determinations of ether extract the outer tube should have an inside diameter of 26 mm, and the inner tube an outside diameter of 22 mm, only 8 to 10 cc. of the solvent being

required. If, however, large amounts of material Fig. 19.—Johnson Extrac(25 to 50 grams) are to be extracted, the diameters tion Tubes.

may be made 32 mm. and 28 mm. respectively and a larger amount of solvent employed.

Where only a few extractions are made, the heating can be performed over (but not on) a metal plate heated by a Bunsen burner, and the condensation effected by an ordinary Liebig condenser. If, however, a considerable number of extractions are carried out, the set apparatus shown in Fig. 20 will be found convenient and also economical of space. It may be attached to the wall or placed at the back of a working desk. The heating, as shown in the cut, is effected by means of two steam pipes, but some form of electric heater answers equally well. The case with glazed door prevents the radiation of heat. At the top is shown the multiple condenser consisting

of a copper tank with block tin tubes. Water is introduced at the left and carried off at the right.

The solvent is best poured through the material, thus obviating in large degree the crawling of the extract. The door should be opened several times during the extraction and kept open for a few minutes for the pur-

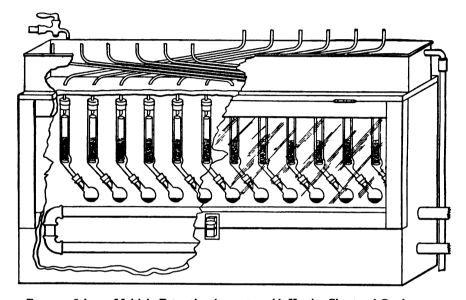


Fig. 20.—Johnson Multiple Extraction Apparatus with Heating Closet and Condenser. pose of rinsing down the sides of the tubes by means of the condensed vapors.

Preparation of Solvents.—In taking the so-called ether extract, sometimes reckoned as fat, the solvent employed is either ethyl ether or the cheaper petroleum ether. Whichever reagent is employed, certain precautions are necessary for the purity of the reagent. If ethyl ether is used, it should be entirely freed from moisture and alcohol by first shaking with water to remove the larger portion of the alcohol, allowing it to stand for some time over dry calcium chloride, and then distilling over metallic sodium. The ether thus prepared should be kept till used with sodium in the container, the latter being somewhat loosely corked, to allow escape of the hydrogen formed.

Petroleum ether is variously termed benzine, naphtha, or gasoline. It should be low-boiling, preferably between 35° and 50°, and it is always best to redistil it before using, in order to be sure it is free from residue. As to the choice of the two reagents for use in fat extraction, it may be said that ethyl ether is the solvent most used, but if a large number of determinations are to be made, the lower cost of petroleum ether is to

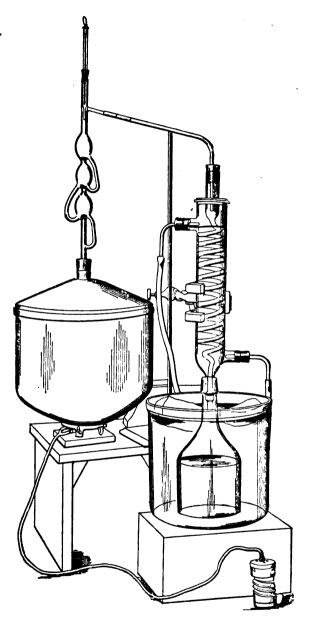


Fig. 21.—Fractionating-still, Arranged for Petroleum Ether.

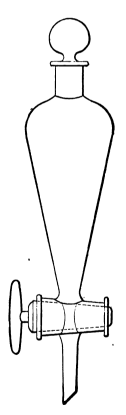


Fig. 22.—A Convenient Form of Separatory Funnel.

be considered. A convenient still for fractionating such substances as petroleum ether is shown in Fig. 21.

Extraction with Immissible Solvents.—It is frequently necessary to dissolve out a substance from a liquid by shaking it with an immiscible solvent, as, for example, in the extraction of certain preservatives from aqueous or acid solutions with ether, petroleum ether, or chloroform. This can be done by shaking in ordinary flasks, but is attended by some difficulty and loss on decantation. A separatory funnel of the type shown in Fig. 22 is almost indispensible for this kind of extraction. The liquid

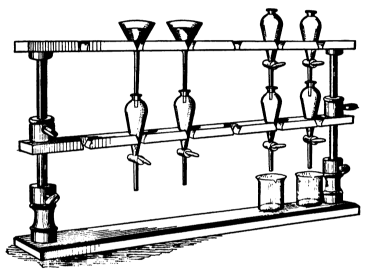


Fig. 23.—Separatory Funnel Support.

and solvent are transferred to the funnel, which is then stoppered and shaken. If the solvent is heavier than water, as in the case of chloroform, it is drawn off from beneath through the outlet-tube of the funnel, closing the tap when the line of demarkation between the two liquids reaches the tap. Or, if the solvent is the lighter, as in the case of ether, the aqueous liquid lying beneath is first drawn off and finally the solvent is poured out through the top. If troublesome emulsions form when shaken, they may frequently be broken up by adding an excess of the solvent and again very gently shaking, or by careful manipulation with a stirring rod. If the solvent is ether, and an obstinate emulsion forms, it may frequently be broken by the addition of chloroform. Such a mixture of ether and chloroform sinks to the bottom and may be drawn off as in the case of chloroform

alone. Ether or chloroform emulsions that refuse to yield to either of the above methods may often be broken by the aid of a centrifuge. A convenient form of separatory funnel support is shown in Fig. 23. It serves for holding the separatory funnels while drawing from one into another, and is also useful as a support for ordinary funnels. The two shelves are adjustable by means of thumb screws. The holes in these shelves are somewhat wider than the slots, so that the separatory funnels after being introduced through the latter drop into position and are held firmly while manipulating the stop-cock.

Determination of Nitrogen by Moist Combustion.—In thus determining nitrogen, the organic matter is first decomposed by digestion with sulphuric acid and an oxidizer, the carbon and hydrogen being driven off as carbon dioxide and water respectively, while the nitrogen is converted into an ammonium salt, from which free ammonia (NH₃) is later liberated by making alkaline. The ammonia is then distilled into an acid solution of known value and calculated by titrating the excess of acid.

In the Kjeldahl process the oxidation is effected by means of a mercury compound, in the Gunning method, by potassium sulphate which forms the bisulphate with the acid.

Neither method in its simplest form is applicable in the presence of nitrates; if these are present, a modification must be used. The Gunning-Arnold method (page 432) is employed for the determination of nitrogen in pepper, as the piperin is not completely decomposed by the usual processes.

The Gunning Method.—Reagents:

Standard alkali solution, N/10 NaOH.*
Pulverized potassium sulphate.
Sulphuric acid, concentrated.
Sodium hydroxide, saturated solution.
Standard acid solution, N/10 H₂SO₄ or HCl.*
An indicator, cochineal.
Granulated zinc.

^{*}Winton employs standard acid of such a strength that I cc. is equivalent to 1% of nitrogen, working on a gram of material, and titrates back with standard alkali two and one-half times weaker than the acid. In order to insure accurate readings, burettes of narrow bore (I cc.=2.6 cm.) are employed. The alkali burette is so graduated that a reading of I corresponds to 2.5 cc., thus allowing for the greater dilution. The advantage of this system is that the per cent of nitrogen is obtained by simply subtracting the alkali reading from the number of cc. of acid employed.

The digestion and distillation are preferably carried out in the same flask, which should be pear-shaped with flat or round bottom and made of moderately thick Jena glass. A convenient size has the following dimensions: length 29 cm., maximum diameter 10 cm., tapering gradually to a long neck, which near the end is 28 mm. in diameter with a flaring edge. Its capacity is about 550 cc.

If desired, the digestion may be conducted in a smaller hard-glass flask of about 250 cc. capacity and of the same shape as the above, and the distillation in an ordinary round-bottomed flask of 500 cc. capacity.

Introduce from 0.5 to 3.5 grams of the sample into the digestion-flask with 10 grams of potassium sulphate and from 15 to 25 cc. of concentrated sulphuric acid. The flask is inclined over the flame and heated gently for a few minutes below the boiling-point of the acid till the frothing has ceased, after which the heat is gradually increased till the acid boils, and the boiling is continued till the contents have become practically colorless or at least of a pale straw color. Wire gauze may be interposed between the flask and flame, but a triangle or a similiar support is to be preferred.

The contents of the flask are then cooled, and, if the digestion has been conducted in the larger flask suitable also for distilling, as above recommended, 300 cc. of water are added and sufficient strong sodium hydroxide to make the contents strongly alkaline, using phenolphthalein as an indicator. If a separate flask is used for the distillation, the contents of the digestion-flask are transferred thereto with the water and the alkali added. A few pieces of granulated zinc should also be introduced, which by the evolution of gas prevents bumping and the sucking back of the distillate. The flask is then well shaken and connected with the condenser, the bottom of which is provided with an adapter, dipping below the surface of the standard hydrochloric or sulphuric acid, a measured quantity of which is contained in the receiving-flask. The distillation is then continued till all the ammonia has passed over into the acid, this part of the operation requiring from forty-five minutes to an hour and a half. As a rule the first 250 cc. of the distillate will contain all the ammonia.

The excess of acid in the receiving-flask is then titrated with standard alkali, and the amount of nitrogen absorbed as ammonia is calculated. The reagents, unless known to be absolutely pure and free from nitrates and

ammonium salts, should be tested by conducting a blank experiment with sugar, by means of which any nitrates present are reduced. Any nitrogen due to impurities should be corrected for.

In purchasing sulphuric acid for nitrogen determination it is important to specify that it be "nitrogen-free" as the so-called chemically pure acid often contains a considerable amount of nitrogen.

Modification of Gunning Method to include Nitrogen of Nitrates.—In addition to the reagents used in the simpler Gunning method, sodium thiosulphate and salicylic acid are required.

A mixture of salicylic and sulphuric acids is made in the proportion of 30 cc. of concentrated sulphuric to 1 gram of salicylic. From 30 to 35 cc. of

Fig. 24.—Bank of Stills for Nitrogen Determination by Gunning Process.

the mixture are added to the 0.5 to 3.5 grams of the substance in the digestion-flask, the flask is well shaken and allowed to stand a few minutes,

occasionally shaking. Then 5 grams of sodium thiosulphate are added, and 10 grams of potassium sulphate, after which the heat is applied, at first very gently and afterwards increasing slowly till the frothing has ceased. The heating is then continued till the contents have been boiled practically colorless. From this point on, proceed as in the Gunning method.

The Kieldahl Method.—One gram of the air dry substance, or a proportionately larger amount of a moist or liquid substance, and 0.7 gram of mercuric oxide (or an equivalent amount of metallic mercury) are placed

Fig. 25.—Johnson Digestion Stand for Nitrogen Determination with Lead Pipe for Carrying off Fumes.

in a 550 cc. Jena flask and 20 cc. of sulphuric acid added. The flask is placed in an inclined position over a Bunsen burner, and the mixture heated below boiling for 5 to 15 minutes or until the frothing ceases, after which the heat is raised until the mixture boils briskly. The boiling is continued until the liquid has become nearly colorless and for a half hour in addition. The lamp is then turned out, the flask placed in an upright position, and potassium permanganate slowly added with shaking until the solution takes on a permanent green or purple color.

After cooling, 250 cc. of water are added, then 25 cc. of potassium sulphide solution (40 grams of the commercial salt in 1 liter of water),

sufficient saturated sodium hydroxide solution to render the solution alkaline, and finally a few grains of granulated zinc, shaking the flask after each addition. Without delay connect with the distillation apparatus, and proceed as in the Gunning method.

Apparatus for Nitrogen Determination.—A bank of stills used by the author in nitrogen determination and in other processes is shown in Fig. 24.

The digestion apparatus shown in Fig. 25 is that devised by Johnson, Winton, and Boltwood. The stand is of cast iron, with holes provided

Fig. 26.—Johnson Distilling Apparatus for Nitrogen Determination.

with three projections that support the flask. The lead pipe with holes for receiving the ends of the flasks serves to carry off the acid fumes.

The Johnson distilling apparatus with accessories by Winton is shown in Fig. 26. The distillation tubes, except for the glass traps and bulb receiver tubes, are of block tin, and are cooled in a copper tank filled with water. The receivers for the distillate are ordinary pint milk bottles.

At the left are two bottles with suspended tubes for measuring the potassium sulphide and sodium hydroxide solutions.

Determination of Ammonia.—A weighed quantity of the finely divided sample, treated with ammonia-free water and made alkaline with magnesium oxide free from carbonate, is distilled into a measured quantity of standard acid (tenth-normal hydrochloric or sulphuric acid) and the amount of ammonia determined by titration.

Determination of Amido-nitrogen.*—In the absence of ammonia, or after the removal of the ammonia as described in the preceding section, the sample is boiled for an hour with 5% hydrochloric or sulphuric acid, which converts the amido-compounds into ammonium salts (chloride or sulphate). Assuming asparagin to be the amido-compound acted upon, the reaction is as follows:

$$2C_4H_8N_2O_3+H_2SO_4+2H_2O=(NH_4)_2SO_4+2C_4H_7NO_4. \\ \begin{array}{c} Asparagin \\ Asparagin \\ \end{array} \\ Asparagin \\ Asparatic acid \\ \end{array}$$

Exactly neutralize the free acid with sodium carbonate, add magnesia (free from carbonate), and distil into standard tenth-normal acid. The ammonia is determined by titration in the usual manner, and its nitrogen represents half of the nitrogen contained in the amido-compound, which it is customary to calculate as asparagin.

Determination of the Various Carbohydrates.—Under title of "Cereals" in Chapter X are given in detail methods for separation and determination of sugar, dextrin, crude fiber, etc.

POISONED FOODS.—Such metallic impurities as are present in food products incidental to their preparation, or as adulterants, will be considered under title of the foods liable to such adulteration.

The detection of highly toxic substances in food, such as arsenic, mercuric chloride, the alkaloids, and other organic poisons that do not occur in food naturally or accidentally, and are present, not as adulterants properly so called, but have been added with criminal intent to do injury, come within the province of the medico-legal chemist or toxicologist rather than that of the food analyst, and are beyond the scope of the present work. The methods involved are similar to those used in the detection of these poisons in the stomach, viscera, and other organs and tissues. The reader is referred in this connection to such treatises as those of Blyth † and Dragendorf. ‡ The analyst is, however, so often called upon to test foods for arsenic that an exception in this case will be made.

^{*} Wiley, Agricultural Analysis, Vol. III. p. 424.

[†] Poisons, their Effects and Detection. London, Griffen & Co., 1906.

[‡] Gerichtlich-chemische Ermittelung von Giften. St. Petersburg, 1876.

Detection of Arsenic.—In testing most food substances for arsenic, it is usually unnecessary to entirely destroy the organic matter, but whenever possible the substance under examination should, by treatment with concentrated nitric and sulphuric acids, be brought into the form of a dry char, which may readily be divided finely by the action of a pestle in a mortar. In this condition the arsenic, which by the process has been oxidized to arsenic acid, may be completely dissolved by continual treatment with boiling water. The hot-water solution containing the extract of the powdered char is then cooled, filtered, and submitted to the Marsh apparatus.

For preliminary treatment of liquids or semi-liquid substances, proceed as directed under arsenic in beer, page 728,

In treating substances like meats, vegetables, and the like, follow in general the directions of Chittenden and Donaldson* for organic tissues, the proportions of acid, etc., being varied to suit special conditions. Heat in a porcelain dish 100 grams of the finely divided substances with 23 cc. of pure concentrated nitric acid at a temperature between 150° and 160° C., stirring occasionally with a glass rod. After the substance has taken on a deep yellow or orange color, remove the dish from the heat, add 3 cc. of pure concentrated sulphuric acid, and stir while the nitrous fumes are given off. The operator should wear a rubber glove to protect the hands. Again heat to about 180° and add while hot, drop by drop, 8 cc. of pure concentrated nitric acid, stirring during the addition of the acid. Then heat at 200° till sulphuric acid fumes come off and a dry carbonaceous mass remains.

This is then pulverized and exhausted with boiling water, and the aqueous solution, when cold, submitted to the Marsh test.

The Marsh Apparatus and its Operation.—Fig. 27 shows a simple form of Marsh apparatus applicable for this work. The generator is provided with a doubly perforated rubber stopper, containing the usual delivery-tube and the entrance-tube. The latter has for convenience a 60° funnel at the top, into which the filter-paper can be folded and the solution containing the extract filtered directly into the generator. A chloride of calcium drying-tube is interposed between the generator and the capillary tube, the latter being drawn from hard arsenic-free tubing.

The apparatus is operated in the usual manner, using arsenic-free

^{*} American Chem. Journal, II. No. 4; Chem. News, Jan. 1881, p. 21.

granulated zinc and dilute sulphuric acid. After running the current of hydrogen long enough through the heated tube to prove the absence of arsenic in the reagents or apparatus, the aqueous solution of the char is poured into the moistened filter at the top of the funnel-tube and allowed to filter slowly into the generator. The length of time necessary to deposit in the capillary tube all the arsenic in the sample, or to prove the absence of arsenic, varies with the conditions, but in general if no darkening of

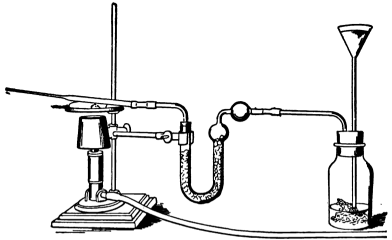


Fig. 27.—Marsh Apparatus for Arsenic.

the tube occurs after an hour, the sample may be considered free from arsenic.

Estimation of Arsenic.*—With the aid of an assay balance sensitive to 0.00001 gram, it is possible to weigh with accuracy an arsenic mirror in a capillary tube when the metallic arsenic amounts to 0.0001 gram or more. Experience will soon show by the appearance of the mirror to the eye when that amount is exceeded. In this case the capillary tube containing the mirror is cut off from the bulk of the tube, and, after drying in a desiccator, is weighed on the assay balance. The capillary is then immersed in a solution of hypochlorite of sodium, which at once dissolves the arsenic only, if present, showing at the same time that the mirror is made up of arsenic and not antimony, which of course would not dissolve. The capillary is then washed, first by water by means of the washbottle, then with a few drops of alcohol, and is finally dried by heat. It

^{*} Leach, Annual Rep. Mass. State Board of Health, 1900, p. 700. Analyst's Reprint, p. 83.

is then cooled and again weighed on the assay balance, the difference in weight corresponding to the metallic arsenic.

If the amount of arsenic is small, it may be estimated by Sanger's method,* which consists in comparing the mirror in the capillary with a series of standard mirrors, made by using varying measured amounts of a standard arsenious oxide solution. This solution is prepared by dissolving I gram of pure arsenious oxide (As₂O₃) in water with the aid of arsenic-free sodium carbonate, and, after acidification with dilute sulphuric acid, making up to a liter. Ten cc. of this solution are measured out carefully and made up to a liter with water, the strength of the dilute solution being 0.01 mgr. to 1 cc. One cc., 2 cc., 3 cc., etc., of this second or dilute solution are separately measured into the Marsh apparatus to give mirrors corresponding to the same number of hundredth-milligrams.

Colorometric Analysis.—Certain analytical processes depend on the formation of a compound of the substance to be determined having a definite color, and the calculation of the quantity present from the intensity of the color of the solution, compared with that of a solution containing a known amount. The comparisons may be made in a special form of cylinder or in a colorimeter. The latter has the advantage that a single solution of known strength serves within reasonable limits for matching any shade in the unknown solution, and for any number of determinations, the desired depth of the color being secured by varying the length of the column.

Schreiner's Colorimeter.†—This apparatus, shown in Fig. 28, consists of two graduated tubes (B), containing the standard and unknown colorimetric solutions, the height of the column of liquid in both tubes being changed by two immersion tubes (A), which remain stationary while the graduated tubes are raised or lowered in the clamps (C). The mirror D reflects the light through the tubes, and the mirror E reflects it again to the eye of the operator at F.

In making the comparisons, the tube containing the solution of either known or unknown strength is set at a definite point, and the other tube is raised or lowered until the colors match. If R is the reading of the standard solution of the strength S, and r the reading of the colorometric solution of unknown strength s, then

$$s = \frac{R}{r}S$$
.

^{*} Proc. Acad. of Arts and Sciences, XXVI. (1891) p. 24.

[†] Jour. Am. Chem. Soc. 27, 1905, p. 1192.

If desired, standard slides of colored glass, such as accompany the Lovibond tintometer, may be used at G for matching the solution of un-

known strength, the value of these slides being determined by comparison with a standard solution.

The Lovibond Tintometer may be used for colorometric chemical analysis, but is not so well suited for this purpose as the Schreiner colorimeter. It is especially designed for determining the color value of liquid and solid technical products, such as beer, wine, oil, flour, paper, etc.

The instrument itself is of simple construction, consisting of an elongated box with an eyepiece at one end and two rectangular openings at the other, one for the solution or substance to be examined, the other for the standard glass slides used for matching the color. Light is reflected through the openings by means of a square piece of opal glass mounted on a jointed standard. Liquids are examined in rectangular cells with glass sides by transmitted light, while powders are pressed into a form and examined by reflected light.

Fig. 28.—Schreiner's Colorimeter with a Tube showing Graduation.

Thus

The standard slides used in general work are red, yellow, and blue in even graduation

from .006 to 20 tint units which can be combined so as to produce any desired tint or shade of any color. The results are expressed in terms of standard dominant colors (red, yellow, and blue), subordinate colors (orange, green, and violet) obtained by combining equal values of two dominant colors, and neutral tint (black) obtained by combining equal values of the three dominant colors.

$$0.6R + 5.6Y = 0.6O + 5.0Y$$

 $0.08R + 1.5Y + 0.2B = 0.08N + 0.12G + 1.3Y$
 $1.2R + 1.0B = 1.0V + 0.2R$

in which R=red, Y=yellow, B=blue, O=orange, G=green, V=violet, N=neutral tint or black.

Special slides may be obtained for the examination of any desired product. For example, slides of brown shades are furnished for beer, of yellow shades for oils, and so on.

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CHAPTER V.

THE MICROSCOPE IN FOOD ANALYSIS.

Microscopical vs. Chemical Analysis.—A very important means of identification of adulterants in many classes of food products is furnished by the microscope, which in many cases affords more actual information as to the purity of food than can be obtained by a chemical analysis. This is especially true of coffee, cocoa, and the spices, wherein the microscope serves to reveal not only the nature of the adulterants, but also not infrequently the approximate amount of foreign matter present. In the case of the cereal and leguminous products so commonly employed as adulterants, a microscopical examination is of paramount importance.

The chemical constants of many of the adulterants of coffee and the spices do not always differ sufficiently from those of the pure foods in which they appear to be distinguished therefrom with accuracy and confidence by a chemical analysis alone. On the other hand, one who is familiar with the appearance under the microscope of the pure foods and of the starches and various ground substances used as adulterants, can, with certainty, identify very minute quantities of these materials, when present, with the same ease that one can recognize megascopically the most familiar objects about him.

A chemical test may, for example, indicate the presence of starch, but it cannot reveal the particular kind of starch. The microscope will at once show whether the starch present is wheat or corn or potato or arrowroot, since these starches differ almost as much in microscopical appearance as do the physical characteristics of the grains or tubers from which they are obtained. Again, by a chemical analysis an abnormal amount of crude fiber may show the presence of a woody adulterant, but only the microscope will enable one to decide whether the impurity consists of sawdust or ground cocoanut shells. Not only in such instances as these is the microscopical examination of greater importance

than a chemical analysis in establishing the purity of the food, but it is at the same time a much quicker guide.

The Technique of Food Microscopy.—The recognition of adulterants by the microscope requires some experience but no more than may be acquired by a chemist who will give the subject serious attention. In the examination of cocoa, coffee, tea, and the spices for adulteration, a careful study of the powdered substance in temporary water mounting will in most cases prove sufficient to familiarize the food analyst with their characteristics under the microscope, and it is not absolutely necessary for him to familiarize himself with the details of section cutting, dissecting, or permanent mounting unless he so desires. The treatment in detail of these latter branches of vegetable histology is beyond the scope of the present work. For full information along these lines the reader is referred especially to such works as those of Behrens*, Zimmerman,† and Chamberlain‡ together with the list of references on page 98.

Standards for Comparison.—For standards the analyst should provide himself with as complete a set as possible of the various materials to be examined, taking care that their absolute purity is established. Whereever possible, he should grind the sample himself from carefully selected whole goods. These, together with samples of the starches and other adulterants, all of known purity, should be contained in small vials carefully stoppered and plainly labeled, arranged alphabetically or in some equally convenient manner in the desk or table on which the microscope is commonly used. The adulterants included in this set of standards should be not only those which experience has shown most liable to be employed, but any which, by reason of their character, might in the analyst's opinion be used under certain conditions.

APPARATUS.

The Microscope-stand.—An expensive or complicated stand is unnecessary. The prime requisites for good work in a microscope-stand are firmness or rigidity, and accuracy in centering. An inexpensive stand possessing these features can be used for the best work, providing the optical parts are satisfactory. It is well, if economy must be practiced, to purchase a simple stand provided with the society screw, and let the larger portion of the allowance go for a high grade of lenses, since many of the attachments inherent in a high-priced stand, though often of convenience, may well be dispensed with.

^{*}Guide to the Microscope in Botany. † Botanical Microtechnique.

† Methods in Plant Histology.

A stand of the so-called continental type (having the horseshoe base) is preferable. A square stage is rather more convenient than the circular form, and the jointed pillar possesses advantages over the rigid variety in ease of manipulation that are certainly worth considering.

The smooth working of both the coarse and fine adjustments should not be lost sight of. If the microscope is to be used exclusively for food work, a substage condenser is unpecessary, hence the construction of the

Fig. 29.—Continental Type of Microscope.

substage may be very simple, unless bacteriological work is to be done as well.

A nose-piece, while not indispensable, is a great convenience for the quick transfer of objectives. A double nose-piece carrying two objectives is ample for routine food work.

The Optical Parts are by far the most important, and should be of superior quality, though not necessarily of the most expensive makers. The food analyst should have at least two objectives, one for high- and one for low-power work, and preferably two oculars.

For the routine examination of powdered food substances the writer prefers a 1-inch objective, used with a medium ocular, the combination giving a magnification of from 240 to 330 diameters, according to the ocular employed. For a low-power objective the 2-inch is a conven-

ient size. It is useful as a finder preliminary to examination with the higher power, and, in connection with a low-power eyepiece, is well adapted for the examination of butter and lard, and for use with the polariscope.

An eyepiece micrometer mounted in an one inch ocular is indispensable for measuring starch grains and other elements. It is calibrated by means of a stage micrometer.

The Micro-polariscope.—This accessory is useful in the identification of starches and other ingredients, and for ascertaining whether or not fats have been crystallized. The polarizer is held below the stage, while the analyzer is applied above the objective, either in the tube or above the ocular.

Fig. 30.—Polarizer and Analyzer for the Microscope.

A common form of construction is one in which the substage is adapted to carry interchangeably the diaphragm tube and the polarizer. If the polariscope is much used, it becomes desirable to provide means for quickly changing the polarizer and diaphragm tube below the stage, and for moving the analyzer in and out of place above the objective. Winton* has devised a microscope-stand with this in view, especially adapted to the needs of the food analyst.

If the polariscope is to be used often, it is convenient to have within easy access two stands, one with the polariscope mounted in place in connection with low-power glasses ready for use, and the other stand trovided with the ordinary high- and low-power objectives only.

Microscope Accessories include of necessity a large number of slides and cover glasses. The latter should be No. 2 thickness, \(\frac{3}{4}\) inch, either round or square.

One or more dissecting-needles in holders and a small hand magnifying-glass should also be provided.

^{*} Journal App. Microscopy, 2, p. 550.

Other useful accessories are a mechanical stage, a pair of fine tweezers, knives, scissors, and, if sections are to be cut, a plano-concave razor.

MICROTECHNIQUE.

Preparation of Vegetable Food Products for Microscopical Examination.—The ground spices and cocoas of commerce are usually of the requisite fineness for direct examination without further treatment. Coffee, chocolate, starches, and similar products should be ground in a mortar fine enough to pass through a sieve with from 60 to 80 meshes to the inch.

A small portion of the powdered sample is taken up on the tip of a clean, dry knife-blade, and placed on the microscope-slide. By means of a medicine-dropper a drop of distilled water is applied, and the wetted

Fro. 31.—Mechanical Stage for Microscope.

powder is then rubbed out under the cover-glass between the thumb and finger to the proper fineness.

The water-mounted slide thus prepared, while useful only for temporary purposes, has proved to be best adapted to the analyst's requirements for routine microscopical examination of powdered food products for adulteration, partly because water is the best medium in most cases for showing up the structural characteristics of these substances and their adulterants, and partly because it serves best for the "rubbing out" process between thumb and finger under the cover-glass, whereby the sample is brought to the requisite degree of fineness.

Experience will soon show how far this rubbing out should be carried for the best effects. Gentle pressure should be applied, care being taken not to break the cover-glass, especially if the sample contain anything of a gritty nature. The rubbing should be continued till the coarser par-

ticles and overlying masses are separated and distributed uniformly, but if too long persisted in, the forms of the tissues, starch grains, and other characteristic portions will be partially destroyed and of too fragmentary a nature to be readily recognizable.

Canada Balsam is the best mountant for the examination of starches under polarized light. In this medium, under ordinary illumination, the starches are not plainly visible, since the refractive index of the balsam is so near that of the starch grains themselves. With the crossed nicols, however, the starch grains stand out very clearly and distinctly in a dark background.

Specimens to be mounted in Canada balsam must be free from moisture. Dehydration is often resorted to by soaking the specimens in alcohol. Canada balsam in solution is prepared by dissolving the balsam broken into small pieces or powdered in a mortar in an equal volume of xylol, filtering and evaporating to sirupy consistency at room temperature.

Glycerin Jelly.*—This is the best permanent mountant for powdered food substances and is prepared as follows: 1 part by weight of the finest French gelatin is soaked two hours in 6 parts of distilled water, after which 7 parts by weight of C. P. glycerin are added, and to each 100 parts of the mixture add 1 part of concentrated carbolic acid. Heat the mixture while stirring till flocculency disappears and filter through asbestos while warm, the asbestos being previously washed and put into the funnel while wet. The jelly is solid at ordinary temperatures, and must be warmed to melt. A small bit of this jelly is removed from the mass by a knife-blade and placed on the cover-glass, which is held over a gas flame till the jelly is melted. The powdered specimen being then shaken into the molten drop, the cover-glass is gently placed upon it (being brought down obliquely to avoid formation of air-bubbles) and pressed down in place.

Microscopical Diagnosis.—It is never safe to pass judgment on a spice or other food by the microscopical examination of a single portion. Several slides should be prepared with bits of the powder taken from different parts of the mass, before the character and extent of the adulteration can be safely determined. Care should be taken that the slide, the knife-blade, the water, and the medicine-dropper be perfectly clean and free from contamination with previous specimens.

It should be borne in mind that at best a composite powdered sample

^{*} Botan. Centralbl., Bd. 1, p. 25.

is but a mechanical mixture of various tissues, and that no two portions will show exactly the same composition.

Characteristic Features of Vegetable Foods under the Microscope.— The structural features of a powdered spice, examined microscopically, will be found to differ considerably in appearance from those of a thin, carefully mounted section of the same spice. Instead of the beautiful arrangement of cells and cell contents with the perfect order of various parts as seen in the mounted section, one finds in the powdered sample under the microscope what often appears to be a most confusing mass of fragments of various tissues. For this reason the most striking characteristics seem to vary with different observers, and it is a well-known fact that microscopists differ widely as to conceptions of size, shape, and ordinary appearance, even in the case of certain of the well-known starch grains. It is on this account that, irrespective of the description of others. the analyst should familiarize himself with the microscopical appearance of the foods with which he is dealing, as well as of their adulterants, forming his own standards as to what constitute the recognizable features, from specimens prepared by himself.

In the large variety of ground berries, buds, tubers, barks, etc., from which the spices and condiments are prepared, as well as in the grains, legumes, shells, fruit stones, and other materials forming the most familiar adulterants, the kinds of plant tissues and cell contents which, under the microscope, serve as distinguishing marks or guides for identification are comparatively few in number.

The most common of these varieties of cell tissue and of cell contents to be met with by the food microscopist in his work are briefly the following:

Parenchyma.—This is most abundant and widely distributed, forming as it does the thin-walled, cellular tissue of nearly all vegetable food substances. The walls of parenchyma cells are, as a rule, colorless and transparent. The forms of the cells are varied and are often sufficiently characteristic in themselves to identify the substance under examination.

Sclerenchyma, or stone cells, are the thick-walled woody cells forming the hard part of nut shells, fruit stones, and seed coverings, occurring also in some fruits and barks. These cells are more often colored and of various shapes but almost always irregular, sometimes elongated, as in cocoanut shells and olive stones occasionally nearly rectangular, as in pepper shells, and sometimes polygonal or nearly circular.

In appearance the sclerenchyma cell commonly has a more or less

deep, central or axial rift, from which small fissures extend through the thick walls, somewhat suggestive of the iris. See Fig. 33.

Variously shaped sclerenchyma cells are found in allspice, cassia,

Fig. 32.—Typical Forms of Various Cell Tissues. Longitudinal section through a clove, showing: Pp, two forms of parenchyma; B, bast fibers; g, vascular and sieve tissue; KK', cells with calcium oxalate crystals. (After Vogl.)

pepper, clove stems, nut shells, etc. Stone cells are optically active to polarized light, and between crossed nicols are very conspicuous by their bright appearance.

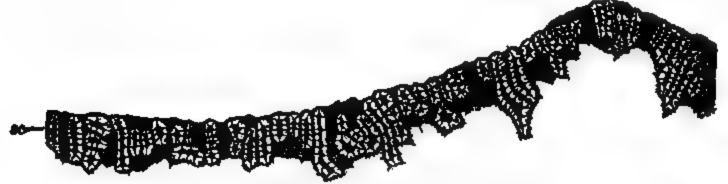


Fig. 33.—Sclerenchyma, or Stone-cell Tissue. A cross-section through the stone-cell layer of the fruit shell of black pepper. (After Vogl.)

Fibro-vascular Bundles are composed of three parts: the bast fibers, or mechanical elements, the phloem, and the xylem.

Bast Fibers are elongated, pointed sclerenchyma cells, of which flax fibers are examples.

Sieve Tubes, the characteristic elements of the phloem, are thinwalled tubes with perforated partitions known as sieve plates.

Vessels or Ducts occur in the xylem. They are designated as spiral, annular, reticulated, or pitted, according to the nature of the walls.

Corky Tissue, or Suberin, constitutes the thin-walled, spongy cells forming the protective, outer dead layers of the bark. This is found in cassia, and in the barks used as adulterants. Suberin is tested for by potassium hydroxide (p. 93).

Starch wherever it occurs furnishes the most characteristic feature of the cell contents, and, as a rule, will at once indicate under the microscope, by the shape and grouping of its granules, the particular substance of which it forms a part. It is very abundantly distributed throughout the vegetable kingdom and occurs in a wide variety of It is particularly conspicuous when viewed by Between crossed nicols such starches polarized light. as corn, potato, and arrowroot show out brightly from a dark background with dark crosses, the bars of which Fig. 34.—Reticulaintersect at the hylum of each granule. When a selenite ory. (After Vogl.) plate is introduced above the polarizer, a beautiful play of colors is seen with various starches, a phenomenon which Blyth applies as a means of identification and classification, but which more modern microscopists regard as of minor importance to distinguishing the various starches morphologically. Starch is found naturally in the cereals, legumes, and many vegetables, in cassia, allspice, nutmeg, pepper, ginger, cocoa, and turmeric. The cereal and leguminous starches from their inertness and cheapness constitute the most common adulterants of the spices and of powdered foods in general. Starch grains are found in the cells of the parenchyma and in other cellular tissues. Iodine is the special reagent (p. 9t).

Gums and Resins occur in characteristic forms among the cell contents of some of the spices. As an example, the portwine-colored lumps of gum in all spice furnish one of the most ready means of recognizing that spice microscopically. Resin is tested for microchemically with alkanna tincture (p. 92).

Aleurone, or Protein Grains, occur in some of the spices, but are not especially characteristic. They somewhat resemble small starch grains. Most varieties of protein grains are soluble in water, but some are insoluble. The soluble varieties, which are not apparent in water-mounted specimens, must be examined in absolute alcohol, glycerin, or oil. In leguminous seeds aleurone occurs closely intermingled with starch in the same cells, while in the cereals it occupies the whole cell.

Protein grains are tested for under the microscope by iodine in potassium iodide, which turns them brown or yellowish brown, and by Millon's reagent, which colors them brick red.

Plant Crystals are not uncommon in the class of substances which the food analyst examines. Among the common forms are the piperin crystals found in pepper. Calcium oxalate occurs in many vegetable products as prismatic crystals, crystal aggregates, or needle-shaped crystals (raphides).

Crystals of calcium carbonate are sometimes met with also, as, for example, in hops. Calcium oxalate crystals are insoluble in acetic acid, while being readily soluble in dilute hydrochloric. Calcium carbonate crystals are soluble with effervescence in both acids. The acid reagents are directly applied to the sample in water-mount under the cover-glass, and the reaction observed through the microscope.

Fat Globules are of common occurrence in many foods and appear of various sizes, sometimes large and conspicuous, and again almost lost sight of because of their minuteness. They are sometimes colorless, as in mace, and sometimes deeply tinted, as in cayenne. Alkanna tincture is used as a reagent for fat (p. 92).

Other Cell Contents of less importance, but which may be identified by the microscope with reagents, are tannic acid (tested for by chloriodide of zinc and ferric acetate (pp. 91 and 92), and various essential oils, for the detection of which alkanna tincture is employed.

REAGENTS IN FOOD MICROSCOPY.

Unless a more extended microscopical investigation of vegetable food substances is contemplated than is involved in the mere identification of adulterants, the analyst will have little need for reagents,* but will depend almost entirely on the form and appearance of the various tissues or tissue fragments, as well as on the abundance, shape, and distribution of such distinctive cell contents as the starches, fat globules, or crystals.

^{*} One reagent that is really necessary on the microscope-table, and will very often be required is iodine in potassium iodide.

Analytical reagents are applied to the water-mounted sample by means of a glass rod or pipette, with which a drop of the reagent is deposited on the sample upon the slide, having previously removed the cover, which is afterwards replaced. Or, without removing the cover-glass, a drop of the reagent is placed in contact with one side of it on the slide. Along the opposite side of the cover is then placed a piece of filter-paper. The latter withdraws by capillary attraction a portion of the water from under the cover-glass, and this is replaced by the reagent, which thus intermingles with the particles of the substance.

Following is a brief list of the commoner microchemical reagents, together with their method of preparation and chief uses. For fuller details in this branch of the subject the reader is referred to Poulsen's Botanical Microchemistry, translated by Trelease, and Zimmerman's Botanical Microtechnique.

A. Analytical Reagents.—Iodine in Potassium Iodide.—Two grams of crystallized potassium iodide are first dissolved in 100 cc. of distilled water and the solution is saturated with iodine.

This reagent is indispensable for the identification of starch, especially when the latter is present in minute quantities. Starch granules when moistened with water are turned blue by iodine, the reaction being exceedingly delicate under the microscope, even when the starch granules are very minute and insignificant without the reagent.

Iodine in connection with sulphuric acid is also useful in distinguishing pure cellulose from its various modifications, such as lignin and suberin. For this purpose the water-mounted sample is first permeated with the iodine reagent, after which concentrated sulphuric acid is applied, with the result that all pure cellulose is turned a deep-blue color, while the modified forms of cellulose are colored yellow or brown. The cellulose is first converted by the sulphuric acid into a carbohydrate isomeric with starch, known as amyloid.

Protein grains are colored brown or yellow brown by the action of iodine.

Chloriodide of Zinc.—Pure zinc is dissolved in concentrated hydrochloric acid to saturation, and an excess of zinc added. The solution is then evaporated to about the consistency of concentrated sulphuric acid, after which it is first saturated with potassium iodide, and finally with iodine.

This reagent may be used instead of sulphuric acid and iodine for the

detection of cellulose, since the zinc chloride converts the cellulose into amyloid, which the reagent colors blue.

Chloriodide of zinc is useful for detecting tannic acid in cell contents. For this purpose the above reagent is much diluted by the addition of a 20% solution of potassium iodide. In this diluted form, when applied to the sample, a reddish or violet coloration is imparted to cell contents having tannin.

Phenol-hydrochloric Acid is prepared by saturating concentrated hydrochloric acid with the purest crystallized carbolic acid. Wood fiber, or lignin, when treated with a drop of this reagent under the cover-glass, and exposed for half a minute to the direct sunlight, will be colored an intense green, which quickly fades.

Indol.—Several crystals of indol are freshly dissolved in warm water. Lignified cell walls assume a deep-red color, when the specimen to be examined is treated first with a drop of the indol reagent, and afterwards washed with dilute sulphuric acid, 1:4.

Millon's Reagent.—This is prepared by dissolving metallic mercury in its weight of concentrated nitric acid, and diluting with an equal volume of water. This reagent, which should be freshly prepared, is of use in testing for protein compounds, which turn brick red when treated with it, especially on gently warming the slide.

Tincture of Alkanna.—A 70 or 80% alcoholic extract of alkanna root, when kept in contact with resins, fixed oils, fats, or essential oils for a short time, stains these cell contents a lively red. The staining is hastened by the aid of heat. Essential oils and resins are soluble in strong alcohol, while fixed oils and fats are insoluble, hence the distinction between these classes of cell contents may be made by the application of alcohol to the alkanna-stained specimen.

Ferric Chloride, Ferric Acetate, or Ferric Sulphate, used in dilute aqueous solution, are all applicable as reagents for tannic acid, which, when present in appreciable amount, will be colored green or blue by either of these reagents.

B. Clarifying Reagents.—Many of the harder cellular tissues are too opaque for careful examination, and may be rendered transparent by clarifying or bleaching. A portion of the powdered sample is either treated with a drop of the reagent under the cover-glass or is allowed to soak for hours or even days in the reagent, using a drop of the same reagent as a medium for examination on the object-glass, instead of water. The clarifying reagents most commonly used are the following:

Chloral Hydrate.—A 60% solution.

Ammonia.—Concentrated, or 28% ammonia is commonly used.

Potassium Hydroxide, used in various degrees of concentration, often in dilute solution, say 5%. This reagent, added to a water mount, causes swelling of the cell wall, and dissolves intercellular substances and protein. It bleaches most of the coloring matters, destroys the starch, and forms soluble soaps with the fats. Potassium hydroxide is also used in testing for suberin, which is extracted from corky tissue on boiling with the reagent, and appears as yellow drops.

Schultze's Macerating Reagent (concentrated nitric acid and chlorate of potassium) is best used by placing the powder or bit of tissue to be treated in a test-tube with an equal volume of potassium chlorate crystals, adding about 2 cc. of concentrated nitric acid, and warming the tube till bubbles are evolved freely, or until the necessary separation of cells is effected. The sample is then removed and washed with water.

By this treatment, bast and wood fibers as well as stone cells are readily separated from other tissues.

Cuprammonia (Schweitzer's Reagent).—This is prepared by adding slowly a solution of copper sulphate to an aqueous solution of sodium hydroxide, forming a precipitate of cupric hydroxide, which is separated by filtration, washed, and dissolved in concentrated ammonia. It should be freshly prepared, and is never fit for use unless it is capable of immediately dissolving cotton. Indeed its chief use is as a test for cellulose, which it readily dissolves. In observing this reaction under the microscope, the powdered specimen under the cover-glass should be only slightly damp before a drop of the fresh reagent is applied. The cell walls are seen to swell up and gradually become more and more indistinct, till they finally disappear.

Cuprammonia is also used as a test for pectose, which occurs in many cell walls, often intermixed with cellulose. When treated with this reagent, cellular tissue containing pectose is acted upon in such a manner that a delicate framework of cupric pectate is sometimes left behind, after the dissolution of the cellulose with which it is mingled.*

PHOTOMICROGRAPHY.

The photomicrograph serves as a simple means of keeping permanent records of unusual forms of adulteration encountered in the course of routine examination. Besides this, the photomicrograph has at times proved its usefulness as a means of evidence in court, showing as it does with faithfulness the presence of a contested adulterant. It is true

^{*} Poulsen, Botanical Micro-chemistry, p. 15.

that from an artistic standpoint the photomicrograph of a powdered sample is often disappointing, due to the fact that ordinarily much of the field is out of focus, unless a very simple homogeneous subject is photographed, as, for instance, starch. As compared with the carefully prepared drawing of a section, which is usually idealized, the photomicrograph is in a sense the more truthful representation.

SUMMARY OF MICROCHEMICAL REACTIONS FOR IDENTIFYING CELLULAR TISSUE AND CELL CONTENTS. BASED ON BEHRENS'.*

		1					
	Iodine in Potassium Iodide.	Chlor- iodide of Zinc.	Iodine and Sul- phuric Acid.	Cupram- monia.	Potassium Hydroxide.	Concen- trated Sulphuric Acid.	Schultze's Mixture.
Cellulose, cell substance.	Yellow to	Violet	Blue	Dissolves	Swells up	Dissolves	Dissolves
Lignin, wood substance.	Yellow	Yellow	Yellow to	Insoluble	Dissolves	Dissolves	Dissolves easily
Middle lamella, inter- cellular substance	Yellow	Yellow	Yellow	Insoluble			Dissolves easily
Suberin, cork substance.	Yellow or brownish	Yellow or brown	Brown	Insoluble	Insoluble in cold. By boiling it comes out in drops	Insoluble	Gives ceric acid reac- tion†
Starch					Dissolves Dissolves		
Protéin	yellow				Dissolves	· · · · · · · · · · · ·	
Gums and resins					.6		
Fat		l			Saponifies		
Tannin		Reddish to violet					
		100 110101			,		1
Calcium oxalate crystals Calcium carbonate							
							Millon's Reagent.
Cellulose, cell substance. Lignin, wood substance.	Phenol- hydro- chloric Acid. Uncolored Green	Indol.	Perric Acetate or Sul- phate.	Alkanna Tincture.	Hydro- chloric	Acetic Acid.	Millon's Reagent.
Cellulose, cell substance. Lignin, wood substance. Middle lamella, intercellular substance	Phenol- hydro- chloric Acid. Uncolored Green	Indol. Uncolored Cherry red Cherry	Perric Acetate or Sul- phate.	Alkanna Tincture.	Hydro- chloric Acid.	Acetic Acid.	Millon's Reagent.
Cellulose, cell substance. Lignin, wood substance. Middle lamella, intercellular substance Suberin, cork substance.	Phenol- hydro- chloric Acid. Uncolored Green Green	Indol. Uncolored Cherry red Cherry red Uncolored	Ferric Acetate or Sul- phate.	Alkanna Tincture.	Hydro- chloric Acid.	Acetic Acid.	Millon's Reagent.
Cellulose, cell substance. Lignin, wood substance. Middle lamella, intercellular substance. Suberin, cork substance.	Phenol- hydro- chloric Acid. Uncolored Green Green Uncolored	Indol. Uncolored Cherry red Cherry red Uncolored	Ferric Acetate or Sul- phate.	Alkanna Tincture.	Hydro- chloric Acid.	Acetic Acid.	Millon's Reagent.
Calcium carbonate Cellulose, cell substance. Lignin, wood substance. Middle lamella, inter- cellular substance. Suberin, cork substance. Starch. Protein. Gums and resins.	Phenol- hydro- chloric Acid. Uncolored Green Uncolored	Indol. Uncolored Cherry red Uncolored	Perric Acetate or Sul- phate.	Alkanna Tincture.	Hydro- chloric Acid.	Acetic Acid.	Millon's Reagent.
Calcium carbonate Cellulose, cell substance. Lignin, wood substance. Middle lamella, intercellular substance. Suberin, cork substance. Suberin, cork substance. Gums and resins. Pat.	Phenol- hydro- chloric Acid. Uncolored Green Uncolored	Indol. Uncolored Cherry red Cherry red Uncolored	Perric Acetate or Sul- phate.	Alkanna Tincture.	Hydro-chloric Acid.	Acetic Acid.	Millon's Reagent.
Cellulose, cell substance. Lignin, wood substance. Middle lamella, intercellular substance. Suberin, cork substance. Starch. Protein. Gums and resins. Fat. Fat. Essential oils.	Phenol- hydro- chloric Acid. Uncolored Green Uncolored	Indol. Uncolored Cherry red Uncolored	Perric Acetate or Sul- phate.	Alkanna Tincture.	Hydro- chloric Acid.	Acetic Acid.	Millon's Reagent.
Calcium carbonate Cellulose, cell substance. Lignin, wood substance. Middle lamella, intercellular substance. Suberin, cork substance. Starch. Protein. Gums and resins. Pat. Resential oils. Tannin.	Phenol- hydro- chloric Acid. Uncolored Green Uncolored	Indol. Uncolored Cherry red Cherry red Uncolored	Perric Acetate or Sulphate.	Alkanna Tincture.	Hydro-chloric Acid.	Acetic Acid.	Millon's Reagent.
Cellulose, cell substance. Lignin, wood substance. Middle lamella, intercellular substance. Suberin, cork substance. Starch. Protein. Gums and resins. Fat. Fat. Essential oils.	Phenol- hydro- chloric Acid. Uncolored Green Uncolored	Indol. Uncolored Cherry red Cherry red Uncolored	Perric Acetate or Sulphate.	Alkanna Tincture.	Hydro- chloric Acid.	Acetic Acid.	Millon's Reagent.

^{*} Microscopical Investigation of Vegetable Substances, page 356.
† When treated with the reagent, suberin forms masses of ceric acid, soluble in ether, alcohol, or chloroform.

While the analyst examines microscopically the ordinary powdered spice, for example, he constantly moves with his hand the fine adjustment-screw, bringing into focus different parts of the field successively. This

he does unconsciously, so that he does not realize how far from flat the field actually is till he undertakes to photograph it, when, as a rule, only a small portion is in good focus. It is therefore impossible in one photograph to show successfully many varied forms of tissue or cell contents in the powder, but separate photographs should be made as far as possible with only a little in each. Thus, for example, with a composite subject like powdered cassia bark, it would be very difficult to show starch, stone cells, and bast fibers in one field, all in equally good focus, and, for the best results only, one, or at most two, such varied groups of elements should be shown in one picture.

Appurtenances and Methods of Procedure.—The temporary method of water-mounting employed by the analyst in routine examination presents many difficulties from a photographic point of view. The vibrating motion of the particles is very annoying, and some skill is required in using just the right amount of water, in avoiding air-bubbles, in waiting the requisite amount of time before exposing the plate for the vibratory motion to cease, and, on the other hand, avoiding too long delay, which would result in the evaporation of the water, and the consequent breaking up of the field. In the writer's experience, however, in spite of these difficulties. the water-mounting gives decidedly the clearest results, and, with patience on the part of the operator, it is in many ways the most desirable method of mounting for photographic purposes. It is in fact the method employed in making most of the photomicrographs of powdered specimens that appear in the plates at the end of this volume, though a few were mounted in glycerin jelly, and the starches for the polarized-light pictures in Canada balsam. The sections of tissues shown in the plates were mounted some in glycerin and others in glycerin jelly.

Experience has shown that two degrees of magnification well calculated to bring out the chief characteristics of the spices and their adulterants in a photomicrograph are 125 and 250 diameters. The starches, which are the most common of any one class of adulterants, vary very widely in the size of their granules. With these the larger magnification of 250 has been found satisfactory, while the general appearance of the composite ground-spice itself under the microscope, as well as that of such adulterants as ground bark, sawdust, chicory, pea hulls, and the like, is best shown with the lower power of 125.*

^{*}The degrees of magnification adopted in the originals of most of the photomicrographs illustrated in the accompanying plates are accordingly 125 and 250, but in the process of lithographing, the photographs were slightly reduced, so that the actual scales in the reproduction are 110 and 220 respectively.

The object, mounted in the manner above described, is best examined when held in a mechanical stage, furnished with micrometer adjustments, in such a manner that a typical field may be selected and held in place long enough to photograph.

The Camera.—This need not of necessity be complicated, but may consist simply of a horizontal wooden base on which the microscope

Fig 35a.—A Convenient Photomicrographic Camera.

rests, and an upright board firmly secured to the base, carrying a frame for an interchangeable ground glass and plate-holder, with a rubber gauze skirt hanging from the frame, adapted to be gathered and tied about the top of the microscope-tube. Means are further provided, as by a slotted guide and screw, for adjusting the frame at any desired height on the upright board.*

A more convenient form of apparatus now employed by the writer is that shown in Figs. 35a and 35b.

^{*} Such a contrivance as this was employed in making some of the accompanying photomicrographs.

The base is a solid iron plate upon which the microscope rests (any microscope may be used with this camera), and above which the camera bellows is supported on two solid steel rods. The bellows is supported at either end on wooden frames.

The ground glass is provided with a central transparent area, formed by cementing a cover-glass upon the ground glass, and permits the accurate focusing of the most delicate detail by means of a hand magnifying-glass. The vertical rods supporting the bellows are attached to metal arms, immovably fixed to a horizontal axis, thus permitting the camera to be tilted

Fig 35b.-Photomicrographic Camera in Horizontal Position

to any angle from vertical to horizontal. It is fixed at the desired angle by means of heavy hand-clamps.

In use the camera is placed in a vertical position and the microscope adjusted on the base so that its tube will coincide with the opening in the front of the camera. The connection between microscope and camera is made light-tight by means of a double chamber, which permits considerable vertical motion of the tube of the microscope without readjustment. A jointed telescoping rod is attached to the upper end of the camera to act as a support, giving perfect steadiness when in a horizontal position, and folding down parallel with the bellows so as to be out of the way when in any other position.

Amplification.—The vertical rods are graduated in inches for determining the amount of amplification, and to show when the ground glass is at right angles to the optical axis. The following simple rule for determining the amount of amplification will give sufficiently accurate results. When photographing without the eyepiece, divide the distance of the ground glass from the stage of the microscope in inches, by the focal length in inches of the objective used. When photographing with the eyepiece, proceed as above and multiply the result by the quotient obtained by dividing 10 by the focus in inches of the eyepiece used.

Adjustment and Manipulation.—The microscope can be placed in any position desired, and the camera adjusted to it. The bellows can then be raised and the microscope used as though no camera were present. When an object is to be photographed, the bellows may be slid into position without in any way disturbing the arrangement of light or object, the final focusing on the ground glass being effected quickly by means of the fine adjustment-screw of the microscope. The exposure having been made, observation through the microscope may be continued without interruption by simply raising the bellows again.

When a water-mounted specimen is to be photographed, the camera and microscope tube should be vertical instead of inclined as shown in the cut.

The camera is best kept in a dark room where the exposures are to be made, the source of light being a 16- or 32-candle-power electric lamp, preferably provided with a ground-glass bulb. The light from this lamp is first carefully centered by moving the reflector of the microscope.

In making pictures, for instance, of the magnification of 250 diameters, the objective, having an equivalent focus of $\frac{1}{6}$ inch, may be used in combination with the one-inch ocular, with the ordinary tube length of microscope. For a lower power, such as 125 diameters, the same objective is employed, but the eyepiece is left out, it being found necessary in this case to remove the upper tube of the microscope, which ordinarily carries the eyepiece, as otherwise the size of the field to be photographed would be restricted. In each case a diaphragm is used in the microscope stage, having an opening of about the same size as that of the front lens of the objective. By means of a stage micrometer scale, the proper position of the camera back is previously determined to give the required magnification.

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CHAPTER VI.

THE REFRACTOMETER.

THE refractive index ranks in importance with the specific gravity as a means of determining the identity and purity of various food products, as well as of estimating the percentage of valuable constituents. Various forms of refractometer are used in food analysis.

The Abbé refractometer is of general application in determining the refractive index of fats, fatty oils, waxes, and essential oils, in estimating the solids in saccharine substances, and in other analytical operations. It employs but a few drops of the material, and reads the refractive index directly, using ordinary white light.

The immersion refractometer, an instrument of recent introduction, is suited for the examination of milk serum to detect added water therein, the detection and determination of methyl alcohol in ethyl alcohol, and the standardization of a wide variety of solutions. The instrument is immersed directly in the liquid to be examined, the degree of refraction being indicated on an arbitrary scale.

The *Pulfrich* is used with the sodium light, and requires a larger amount of material than the Abbé, the liquid being held in a cylinder above the prism. The scale reading is in angular degrees, from which the index of refraction is calculated by a formula or from a table. The instrument is provided with a temperature-controlling apparatus.

In the Amagat and Jean or oleo-refractometer, an outer and an inner cylinder are respectively filled with an oil of known value or purity, and with the oil to be examined. By the comparative displacement to the right or left of a beam of white light passing through both cylinders, the displacement being read in degrees on an arbitrary scale, the refraction of an oil may be measured. Two oils may thus be readily compared under the same conditions, one of known purity, for example, with a doubtful sample of the same kind.

The butyro-refractometer and the Wollny milk fat refractometer are, as their names imply, instruments primarily intended for more restricted fields of work than the foregoing. They involve the same principle as the Abbé, but are simpler in construction and have arbitrary scales.

Unless such widely varying substances as the waxes and the essential oils are to be studied, the Zeiss butyro-refractometer, though primarily

intended for the examination of butter and lard, answers most of the purposes of the Abbé instrument with the advantage of greater simplicity, being equally well adapted for examining all the common edible oils and fats, as well as other materials.

THE ZEISS BUTYRO-REFRACTOMETER.

This instrument (shown in Fig. 36) is so constructed that the degree of refraction of a beam of light, which passes obliquely through a thin

Fig. 36.—The Zeiss Butyro-refractometer.

film of the fat, is read on an arbitrary scale of sufficient extent to cover the widest limits of deviation possible for butter fat and oleomargarine under ordinary temperatures.

The graduation is in divisions from 1 to 100, covering a variation in refractive indices of from 1.4220 to 1.4895. A and B are the two hinged hollow portions of the prism casing of the instrument, so arranged that when closed together the melted fat is held in a film of sufficient thickness between the two opposed transparent prism surfaces, the beam of light, either diffused daylight or lamplight, being reflected through it by means of the mirror J. The transparent scale is within the telescope tube at the height indicated by G.

The refractometer is connected to any kind of heating arrangement. which admits of warm water being transmitted through the prism casing. in at D and out at E. A simple arrangement, which suffices for all ordinary cases, may expeditiously be improvised in the following manner: Fill a vessel of say 2 gallons capacity with water of 40° to 50° C. Into this vessel dip the free end of an india-rubber tube slipped over the nozzle D and let the vessel be placed at a height of about one-half or one vard above the refactometer. Then it will be seen that suction at a tube attached to E will cause the warm water to flow through the prism casing by the action of the siphon arrangement. By means of a pinch clip the velocity of the water may be regulated at will. The waste water may be allowed to flow into a second vessel and, provided its temperature does not fall below 30°, it may be used for replenishing the upper vessel.

When working with solid fats, a temperature must be maintained by the heated water well above the melting-point of the fat. With liquid oils no heater is necessary, as determinations may be made at room temperature, but it is advisable in all cases to have a constant stream of water passing through the water jacket, which may be done by directly connecting it with the water faucet in the case of oils, since, without such precautions to insure even temperature, disturbing variations are liable to occur, due to the warming of the prisms by the manipulation of cleaning, etc.

Refractometer Heater.—A regular heater, shown in Fig. 37, is furnished by the manufacturers, capable of supplying a current of water of approximately constant temperature, and will be found of great convenience when the instrument is to be used constantly, especially with the solid fats.

A supply reservoir A is secured to the wall and is connected by means of a rubber inlet tube G to the water faucet C. The reservoir is provided with a waste overflow pipe and with an outlet tube D, the flow through the latter being regulated by the cock H. The tube D leads to the spiral heater HS, which is heated by a Bunsen burner. From the heater the tube E conducts the warm water through the refractometer, from which it flows through the tube F, either directly into the sink, or into the intermediate vessel B. The temperature of the water is regulated by adjusting the cock H, and the height of the flame of the Bunsen burner.

Manipulation of the Butyro-refractometer.—The prism casing is first opened by giving about half a turn to the right to the pin F, Fig. 36, until it meets with a stop. Then simply turn the half B of the prism

casing aside. Pillar H holds B in the position shown in Fig. 36. The prism and metallic surfaces must now be cleaned with the greatest care, the best means for this purpose being soft linen, moistened with a little alcohol or benzine.

If the sample to be examined is a solid fat, maintain the temperature above the melting-point, and apply by a glass rod a drop or two of the clear melted fat (filtered if turbid) to the surface of the prism contained in the casing B. For this purpose the apparatus should be raised with

Fro. 37.—The Zeiss Heating Apparatus for all Forms of Refractometer. Shown in the cut in connection with the Pulfrich refractometer.

the left hand so as to place the prism surface in a horizontal position. A liquid oil is directly applied in the same manner without preliminary precautions as to heating. Now press B against A, and place F by turning it in the opposite direction, in its original position; thereby B is prevented from falling back, and both prism surfaces are kept in close contact. Place the instrument again upon its sole plate.

While looking into the telescope, give the mirror J such a position as to render the critical line, which separates the bright left part of the field from the dark right part, distinctly visible. It may also be necessary to move or turn the instrument about a little. First it will be necessary to ascertain whether the space between the prism surfaces be uniformly filled with oil or fat, failing which the critical line will not be distinct. For this purpose examine the rectangular image of the prism surface formed about 1 cm. above the ocular with a hand magnifier or with the

naked eye, placing the latter at its proper distance from the ocular. Finally adjust the movable front part of the ocular so that the scale becomes clearly visible.

By allowing a current of water of constant temperature to flow through the apparatus some time previous to the taking of the reading, the at first somewhat hazy critical line approaches in a short time, generally after a minute, a fixed position, and quickly attains its greatest distinctness. When this point has been reached, note the appearance of the critical line (i.e., whether colorless or colored, and in the latter case of what color); also note the position of the critical line on the centesimal scale, which admits of the tenth divisions being conveniently estimated; at the same time read the position of the thermometer.

Testing the Adjustment of the Ocular Scale.—It is imperative that the adjustment of the instrument be tested periodically, and in particular when it is being used for the first time. This may be done by means of the standard fluid supplied with the instrument, the critical line of which is approximately colorless, and must occupy the following positions in the scale.

Temper-	Scale	Temper-	Scale	Temper-	Scale	Temper-	Scale
ature.	Division.	ature.	Division.	ature.	Division.	ature.	Division.
30°	68.1	25°	71.2	20°	74·3	15° 14° 13° 12° 11° 10°	77-3
29°	68.7	24°	71.8	19°	74·9		77-9
28°	69.3	23°	72.4	18°	75·5		78.6
27°	70.0	22°	73.0	17°	76·1		79-2
26°	70.6	21°	73.6	16°	76·7		79.8
25°	71.2	20°	74.3	15°	77·3		80-4

The fractional parts of a degree can accordingly easily be brought into calculation (0.1=0.06 scale div.). Deviations of 1 to 2 decimals of the scale divisions are of no consequence, and are in most cases due to inexact determinations of temperature. Should, however, careful tests result in the discovery of greater deviations, readjustment of the scale will be necessary, which may be effected by means of a watch-key supplied with the instrument, in accordance with the values given in the above table. The watch-key is inserted at G in Fig. 36, and by its means the position of the objective, and therefore that of the critical line with respect to the scale may be altered.

Transformation of Scale Divisions into Indices of Refraction.—The following table, adapted from that of Pulfrich, enables one to convert scale readings on the butyre-refractometer into indices of refraction, n_D and vice versa:

EQUIVALENTS OF INDICES OF REFRACTION AND BUTYRO-REFRACTOMETER READINGS.

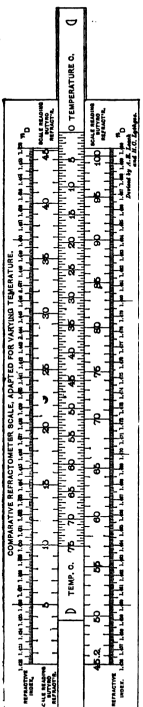
Refrac-		Fourth Decimal of n_{D_i}											
tive Index. *D.	0	1	2	3	4	5	6	7	8	9			
				80	CALE RE	EADINGS	i.						
1.422	0.0	0.1	0.2	0.4	0.5	0.6	0.7	0.9	1.0	1.1			
1.423	1.2	1.4	1.5	1.6	1.7	1.9	2.0	2.1	2.2	2.4			
1.424	2.5	2.6	2.7	2.8	3.0	3.1	3.2	3-3	3-5	3.6			
1.425	3.7	3.8 5.1	4.0	4.I	4.2	4-3	4-5	4.6	4.7 6.0	4.8 6.1			
1.426 1.427	5.0 6.2	6.4	5.2 6.5	5.4 6.6	5·5 6.8	5.6 6.0	5·7 7.0	5.9 7.1	7.2	7.4			
1.428	7.5	7.6	7.7	7.9	8.0	8.1	8.2	8.4	8.5	8.6			
1.429	8.7	8.9	9.0	9. í	9.2	9-4	9.5	9.6	9.8	9.9			
1.430	10.0	10.1.	10.3	10.4	10.5	10.6	10.7	10.9	11.0	11.1			
1.431	11.3	11.4	11.5	11.6	11.8	11.9	12.0	12.2	12.3	12.4			
1.432	12.5 13.8	12.7 14.0	12.8 14.1	12.9	13.0	13.2 14.5	13.3 14.6	13.5 14.7	13.6 14.9	13.7			
1.433	15.1	15.3	15.4	15-5	15.6	15.8	15.9	16.0	16.2	16.3			
1.435	16.4	16.6	16.7	16.8	17.0	17.1	17.2	17.4	17.5	17.6			
1.436	17.8	17.9	18.0	18.2	18.3	18.4	18.5	18.7	18.8	18.9			
1.437	19.1	19.2	19.3	19.5	19.6	19.7	19.8	20.0	20. I	20.3			
1.438	20.4	20.5	20.6	20.8	20.9	21.1	21.2	21.3	21.4	21.6			
1.439	21.7	21.0	22.0	22.1	22.2	22.4	22.5	22.6	22.7	22.9			
1.440	23.0	23.2	23.3	23-4	23.5	23.7	23.8	23.9	24. I	24.2			
1.441	24.3	24.5 25.8	24.6	24.7	24.8	25.0	25.1	25.2	25.4	25.5			
1.442	25.6	25.0 27.1	25.9 27.3	26.1 27.4	26.2 27.5	26.3 27.7	26.5 27.8	26.6 27.9	26.7 28.1	26.9 28.2			
1.443	27.0 28.3	28.5	28.6	28.7	28.9	29.0	29.2	29.3	20.1	29.6			
1.445	29.7	29.9	30.0	30.1	30.3	30.4	30.6	30.7	30.8	30.9			
1.446	31.1	31.2	31.4	31.5	31.6	31.8	31.9	32.1	32.2	32.3			
1.447	32.5	32.6	ე₂.8	32.9	33.0	33-2	33-3	33-5	33.6	33-7			
1.448	33-9	34.0	34.2	34-3	34-4	34.6	34.7	34.9	35.0	35.1			
1.449	35-3	35-4	35.6	35.7	35.8	36.0	36.1	36.3	36.4	36.5			
1.450	36.7	36.8	37.0	37.1	37.2	37-4	37-5	37-7	37.8	37.9			
1.451	38.1	38.2	38.3	38.5	38.6	38.7	38.9	39.0	39.2	39-3			
I.452	39·5 40.9	39.6 41.0	39-7 41.1	39.9	40.0 41.4	40.1	40.3	40.4 41.8	40.6 42.0	40.7 42.I			
1.453	42.3	42.4	42.5	41.3	42.8	43.0	43.I	43-3	43.4	43.6			
1.455	43-7	43-9	44.0	44.2	44.3	44.4	44.6	44.7	44.9	45.0			
1.456	45.2	45.3	45-5	45.6	45.7	45-9	46.0	46.2	46.3	46.4			
1.457	46.6	46.7	46.9	47.0	47.2	47-3 48.8	47.5	47.6	47 - 7	47-9			
1.458	48.0	48.2	48.3	48.5	48.6		48.9	49.1	49.2	49 4			
1.459	49-5	49-7	49.8	50 .0	50.1	50.2	50.4	50 .5	50.7	50.8			
1.460 1.461	51.0 52.5	51.1 52.7	51.3 52.8	51.4 53.0	51.6 53.1	51.7 53.3	51.9 53.4	52.0 53.6	52.2 53-7	52-3 53-9			
1.462	54.0	54.2	54-3	54-5	54.6	54.8	55.0	55.1	55-3	55.4			
1.463	55.6	55.7	55.9	56.0	56.2	56.3	56.5	56.6	56.8	56.9			
1.464	57.1 58.6	57·3 58.8	57·4 58.9	57.6	57-7	57-9	58.0	58.2	58.3	58.5			
1.465	58.6	58.8	58.9	59.1	59.2	59-4	59.5	59-7	59.8	60.0			
1.466	60.2	60.3 61.8	60.5	60.6	60.8	60.9	61.1	61.2	61.4 62.0	61.5			
1.467	61.7 63.2	63.4	62.0 63.5	62.2	62.3 63.8	62.5 64.0	64.2	64.3	64.5	63.1			
									66.1	66.2			
1.469	64.8	65.0	65.1	65.3	65.4	65.6	65.7	65.9	66.1	6			

EQUIVALENTS OF INDICES OF REFRACTION AND BUTYRO-REFRACTOMETER READINGS—(Continued).

Refrac-				Fo	urth Deci	mal of #/	D.			
Index,	0	1	2	3	4	5	6	7	8	9
				80	CALE RI	EADINGS	.			
1.470	66.4	66.5	€6.7	66.8	67.0	67.2	67.3	67.5	67.7	67.8
1.471	68.0	68.1	68.3	68.4	68.6	68.7	68.9	69.1	69.2	69.4
1.472	69.5	69.7	69.9	70.0	70.2	70.3	70.5	70.7	7ó.8	71.0
1.473	71.1	71.3	71.4	71.6	71.8	71.9	72.1	72.2	72.4	72.5
1.474	72.7	72.9	73.0	73.2	73-3	73.5	73-7	73.8	74.0	74-
1.475	74-3	74-5	74.6	74.8	75.0	75.1	75-3	75-5	75.6	75-8
1.476	76.0	76.1	76.3	76.5	76.7	76.8	77.0	77.2	77-3	77-5
1.477	77-7	77-9	78. I	78.2	78.4	78.6	78.7	78.9	79.1	79-2
1.478	79-4	79.6	79.8	80.0	80.1	80.3	80.5	80.6	80.8	81.0
1.479	81.2	81.3	81.5	81.7	81.9	82.0	82.2	82.4	82.5	82.7
1.480	82.9	83.I	83.2	83.4	83.6	83.8	83.9	84.1	84.3	84.
1.481	84.6	84.8	85.0	85.2	85.3	85.5	85.7	85.9	86.o	86.2
1.482	86.4	86.6	86.7	86.9	87.1	87.3	87.5	87.6	87.8	88.0
1.483	88.2	88.3	88.5	88.7	88.9	89.i	89.2	89.4	89.6	89.8
1.484	90.0	90.2	90.3	90.5	90.7	90.9	91.1	91.2	91.4	91.0
1.485	91.8	92.0	92.1	92.3	92.5	92.7	92.9	93.0	93.2	93-4
1.486	93.6	93.8	94.0	94.1	94-3	94-5	94-7	94.8	95.0	95-
1.487	95-4	95.6	95.8	96.0	96.1	96.3	96.6	96.7	96.9	97.0
1.488	97.2	97-4	97.6	97.8	98.0	98.1	98.3	98.5	98.7	98.9
1.489	99.1	99.2	99-4	99.6	99.8	100.0				

The Critical Line.—It should be remembered that the instrument is primarily intended for use with butter, and that the prisms are so constructed that the critical line of pure butter is colorless, while various other fats and oils, notably oleomargarine, which have greater dispersive powers than natural butter, show a colored critical line. When too great dispersion is apparent to render possible an accurate reading, or when the critical line presents very broad fringes, as with linseed oil, poppyseed oil, turpentine, etc., use a sodium light, obtained by the application of table salt to the Bunsen gas flame, or the diffused daylight may be reflected in the mirror through a flat bottle filled with a dilute solution of potassium bichromate, to give a yellow light.

The advantages of the refractometer for examination of fats and oils consist in the convenience with which very accurate determinations of the refractive index may be made at any temperature between 10° and 50° C., inclusive of thermal variations of refractive powers, and also in the possibility which it affords of distinguishing substances by their different dispersive powers, rendered visible by the different coloring of the critical line, a red-colored critical line being indicative of a relatively low dispersive power, a blue line of relatively high dispersion.



Variation of Reading with the Temperature.— No definite temperature has been adopted as a standard for readings of this instrument, but it is easy to reduce readings at any temperature to terms of any other temperature for purposes of comparison. While the change in index of refraction for 1° C. is the same whatever the temperature, as Tolman and Munson have pointed out,* the change in scale reading per 1° C. decreases as the temperature increases, and varies slightly with different oils. For correcting reading R' at a temperature T' to a reading R at temperature T, their formula is R = R' - X(T - T'), X being the change in scale reading due to change of 1° C. in temperature.

For butter, oleomargarine, beef tallow, lard, and other fats reading from 40° to 50° or thereabouts on the scale, X=0.55. For oils reading between 60° and 70° , like olive, mustard, rapeseed, cottonseed, peanut, etc., X=0.58, and for oils reading between 70° and 80° , like corn oil, X=0.62.

The slide rule † shown in Fig. 38, for use with the refractometer, has been jointly devised by H. C. Lythgoe and the writer, to render unnecessary the use of tables or formulas. The extreme upper and lower scale divisions indicate indices of refraction, and adjacent to these are the scale divisions indicating readings on the butyro-refractometer. By comparison, therefore, the values of either the Abbé or the butyro scale may be readily ascertained in terms of the other.

The sliding scale, expressing temperature readings in degrees centigrade, is intended to be used in connection with the adjacent scale of butyro-refractometer readings, to readily express the butyro-scale reading of any fat or oil taken at a given temperature, in terms of that at any other temperature. This is frequently convenient

Fig. 38.—Comparative Refractometer Scale.

^{*} Jour. Am. Chem. Soc., XXIV, p. 755.

[†] Manufactured by Messrs. Baird and Tatlock, Ltd., 14 Cross Street, Hatton Garden, London.

in comparing the work of various observers, where different temperatures have been employed.

The correction for change in n_D on the scale is 0.000365 for 1° C., being based on the experimental work of Tolman, Long, Proctor, Lythgoe, and the author.

THE ABBÉ REFRACTOMETER.

This instrument, Fig. 39, has a much wider range in reading than either the butyro or the Wollny instruments already described, read-

Fig. 39.—The Abbé Refractometer with Temperature-controlled Prisms.

ing as it does to the fourth decimal between the limits of 1.3 and 1.7 in indices of refraction. The equivalent readings of the Wollny milk fat refractometer, in indices of refraction, range from 1.3332 to 1.4220, while those of the butyro instrument run from 1.4220 to 1.4895. The Abbé instrument is thus necessary for use with the high-refracting essential

oils. Its construction is such that the prisms can withstand a higher heat than in the case of the butyro-refractometer, and it is hence better adapted for the examination of samples having a high melting-point, such as beeswax and paraffin. An advantage of the Abbé over the butyro instrument lies in the fact that the wide dispersion, inevitable when reading many substances on the butyro, may be entirely compensated for with the Abbé, and a clear sharp line be obtained. The construction of the prisms in relation to the heating jacket is similar in both instruments, and a film of the substance to be examined is held in the same manner between the surfaces of the prisms.

Construction and Manipulation.—The Abbé refractometer is mainly composed of the following parts (see Fig. 39):

- 1. The double Abbé prism AB, which contains the fluid and can be rotated on a horizontal axis by means of an alidade.
- 2. A telescope OF for observing the border-line of the total reflection which is formed in the prism.
- 3. A sector S, rigidly connected with the telescope, on which divisions representing refractive indices are engraved.

The double prism (AB, Fig. 39) consists of two similar prisms of flint-glass, each cemented into a metal mount and having a refractive index $n_D = 1.75$. The former of the two prisms, that farthest from the telescope, which can be folded up or removed, serves solely for the purpose of illumination, while the border-line is formed in the second flint prism. A few drops of the fluid to be investigated is deposited between the two adjoining inner faces of the prisms in the form of a thin stratum, about 0.15 mm. thick.

The double prism is opened and closed by means of a screw-head v, which acts in the manner of a bayonet catch. In order to apply a small quantity of fluid to the prisms without opening the casing, the screw v is slackened and a few drops of fluid poured into the funnel-shaped aperture of a narrow passage, not seen in the figure. On again tightening the screw, the fluid is distributed by capillary action over the entire space between the two prisms. This arrangement facilitates the investigation of rapidly evaporating fluids, such as ether solutions. In the case of viscous fluids (resins, etc.), a drop of moderate size is applied with a glass rod to the dull prism surface, the double prism being opened for the purpose. The prisms are then closed again, and before the measurement is proceeded with, the refractometer is left standing for a few minutes in order to compensate for any cooling or heating of the prisms which might occur while they were separated.

The arrangement for controlling the temperature of the prisms of the Abbé refractometer is essentially after Dr. R. Wollny's plan of enclosing the prisms in a metal casing with double walls, through which water of a given temperature is circulated.

The border-line is brought within the field of the telescope OF by rotating the double prism by means of the alidade in the following manner: Holding the sector, the alidade is moved from the initial position at which the index points to $n_D = 1.3$, in the ascending scale of the refractive indices until the originally entirely illuminated field of view is encroached upon from the direction of its lower half by a dark portion: the line dividing the bright and the dark half of the field then is the "border-line." When daylight or lamplight is being employed, the border-line, owing to the total reflection and the refraction caused by the second prism, assumes at first the appearance of a band of color, which is quite unsuitable for any exact process of adjustment. conversion of this band of color into a colorless line sharply dividing the bright and dark portions of the field is the work of the compensator, which consists of two similar Amici prisms of direct vision for the D-line, and rotated simultaneously, though in opposite directions, round the axis of the telescope by means of the screw-head M. dispersion of the border-line, which appears in the telescope as a band of color, can thus be counteracted by rotating the screw-head M till the equal but opposite dispersions are neutralized, making the line colorless and sharp.

The border-line is now adjusted upon the point of intersection of the crossed lines by slightly inclining the double prism to the telescope by means of the alidade. The position of the pointer on the graduation of the sector is then read by the aid of the magnifier attached to the alidade. The reading supplies the refractive index n_D of the substance under investigation without any computation, and with a degree of exactness approaching to within about two units of the fourth decimal. Simultaneously, the reading of the scale on the drum of the compensator (T in Fig. 39) enables the mean dispersion to be arrived at by means of a special table and a short process of computation.

Influence of Temperature.—As the refractive index of fluids varies with their temperature, it is of importance to know the temperature of the fluid contained in the double prism during the process of measurement.

If, therefore, it is desired to measure a fluid with the highest degree of accuracy attainable with the Abbé refractometer (to within one or

two units of the fourth decimal of n_D), it is absolutely necessary to bring the fluid, or rather the double prism containing it, to a definite known temperature, and to be able to control this temperature so as to keep it constant to within some tenths of a degree for a period of several hours; hence a refractometer principally required for the investigation of fluids must be provided with heatable prisms.

The type of heater shown in Fig. 37 and described in connection with the butyro-refractometer on page 102, is equally adapted for controlling the temperature of the prisms in the Abbé instrument, the flow of water entering at D and passing out at E, Fig. 39.

THE IMMERSION REFRACTOMETER.

This form of refractometer is of more recent introduction than the others made by Zeiss, and has many features that especially commend it to the use of the food analyst. The construction of the immersion refractometer is such that, as its name implies, it may be immersed directly in an almost endless variety of solutions, the strength of which, within limits, may be determined by the degree of refraction read upon an arbitrary scale. Thus, for example, the strengths of various acids and of a variety of salt solutions used as reagents in the laboratory, as well as of formaldehyde, of sugars in solution, and of alcohol, are all capable of determination by the use of the immersion refractometer.

Figure 40 shows the form used by the writer. P is a glass prism fixed in the lower end of the tube of the instrument, while at the top of the tube is the ocular Oc, and just below this on a level with the vernier screw Z is the scale on which is read the degree of refraction of the liquid in which the prism P is immersed. The tube may be held in the hand and directly dipped in the liquid to be tested, this liquid being contained in a vessel with a translucent bottom, through which the light is reflected. The preferable method of use is, however, that shown in the cut.

A is a metal bath with inlet and outlet tubes, arranged whereby water is kept at a constant level. The water is maintained at a constant temperature by means of a controller of the same type as the refractometer heater shown in Fig. 37. In the bath A are immersed a number of beakers, containing the solutions to be tested. T is a frame on which is hung the refractometer by means of the hook H, at just the right height to permit of the immersion of the prism P in the liquid in any of the beakers in the row beneath. Under this row of beakers the bottom of the tank is composed of a strip of ground glass, through which light is reflected by an adjustable pivoted mirror.

The temperature of the bath is noted by a delicate thermometer immersed therein, capable of reading to tenths of a degree.

Returning to the main refractometer-tube, R is a graduated ring or collar which is connected by a sleeve within the tube with a compound prism near the bottom, the construction being such that by turning the collar R one way or the other the chromatic aberration or dispersion of any liquid may be compensated for, and a clear-cut shadow or critical line projected across the scale. By the graduation on the collar R, the degree of

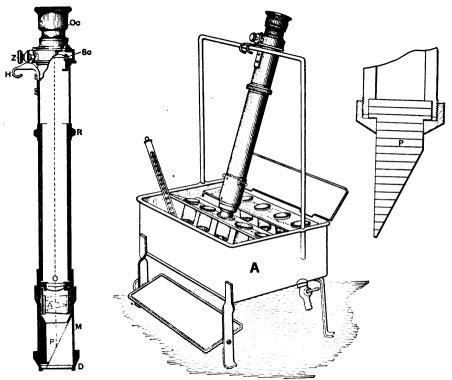


Fig. 40.—The Zeiss Immersion Refractometer.

dispersion may be read. Tenths of a degree on the main scale of the instrument may be read with great accuracy by means of the vernier screw Z, graduated along its circumference, the screw being turned in each case till the critical line on the scale coincides with the nearest whole number.

The scale of the instrument reads from -5 to 105, corresponding to indices of refraction of from 1.32539 to 1.36640. It should be noted that the index of refraction may be read with a greater degree of accuracy on the immersion refractometer than on the Abbé instrument.

Manipulation of the Immersion Refractometer.—Before using the instrument for the first time, it is necessary to see that the refractometer is correctly adjusted. For this purpose the bath A is placed with its long side parallel to the window and the mirror turned towards a bright sky, the bath is half filled with tap-water, and a beaker filled with distilled water is then placed in one of the five holes in the front row immediately above the mirror. Finally, the refractometer is hung by its hook H upon the wire frame, the prism being totally submerged in the water contained in the beaker.

The whole apparatus is now allowed to stand for ten minutes, or until the distilled water has acquired the exact temperature of the bath, and the ocular is focussed upon the divisions of the scale by turning the milled zone of the ocular shell until the lines and numbers are seen quite distinctly, the mirror being adjusted so that the light of the bright sky is seen directly through the beaker. The upper part of the field from -5 to about 15 appears bright, and it is separated from the lower dark part by a sharp line of demarcation, if the index on the ring of the compensator stands at 5.

SCALE READING AND INDEX OF REFRACTION OF DISTILLED WATER AT 10-30° C., ACCORDING TO WAGNER.

Temper- ature C.	Scale Reading.	Index of Refraction, *D.	*D Difference for 1° C.	Temper- ature C.	Scale Reading.	Index of Refraction, *D.	*Difference for 1° C.
30	11.8	1.33196		19	14.7	1.333075	8.5
29 28	12.1	1.33208	12.0	1 18	14.9	1.33316	8.5
28	12.4	1.332195	11.5	17.5	15.0	1.33320	4 } 8.0
27	12.7	1.33231	11.5	17	15.1	1.33324	4 5 8.0
26	13.0	1.33242	11.0	16	15.3	1.333315	7.5
25	13.25	1.332525	10.5	15	15.5	1.33339	7.5
24	13.5	1.332625	10.0	14	15.7	1.33346	7.0
23	13.75	1.33272	9.5	13	15.85	1.333525	6.5
22	14.0	1.33281	9.0	12	16.0	1.33359	6.5
21	14.25	1.33290	9.0	11	16.15	1.33365	6.0
20	14.5	1.33299	9.0	10	16.3	1.333705	5.5

The reading is taken and the temperature of the distilled water noted. Reference to the above table will show if the refractometer is correctly adjusted. Should the average of several careful readings at a given temperature deviate from that contained in the table, the following should be resorted to:

Readjustment of the Scale.—The ocular end of the refractometer hanging on the wire frame is grasped from behind with the thumb and forefinger of the left hand, the micrometer drum set to 10, and the steel

spike, housed in the case of the apparatus, inserted into one of the holes of the nickeled cross-holed screw lying on the inner side of the micrometer drum. The spike is then turned anti-clockwise, as seen from the rear, whereupon the nickeled milled nut, which governs the micrometer, becomes loosened. The temperature of the distilled water in the beaker is taken once more to see that it has remained constant, and then the table (page 113) is consulted to find the "adjusting number" properly belonging to the temperature indicated. By turning the spike, the borderline is brought exactly upon the integer scale division appertaining to the adjusting number, and the loose micrometer drum is turned so that the index accords with the decimal portion of the adjusting number. The drum is now held firmly with the thumb and forefinger of the left hand, while the nut is screwed up tight again by the right hand, taking care, however, that the drum does not wander off the index. Finally, the new adjustment is tested by repeated readings.

Regulating the Temperature.—In many cases it suffices to allow water at the temperature of the room to flow slowly from a tank suspended high upon the wall through the bath. Should it be required, however, to maintain a given temperature (say 20° C.) for hours together constant to a tenth of a degree, which is frequently desirable if not actually necessary, a more elaborate temperature-regulating device should be employed. In cold weather, or when the tap-water has a lower temperature than that desired, a refractometer heater of the type shown in Fig. 37, and described on page 102, is convenient.

When, as in the summer, the tap-water temperature is higher than that desired for the refractometer bath, there are various ways of successfully controlling the temperature at a lower degree. An ice-water tank placed above the level of the bath may be employed, the flow from which through the bath is carefully controlled by a pinch-cock or otherwise, or is allowed to mingle, under careful regulation before entering the bath, with the water from the tap direct or with that from the heater.

Investigation of Solutions in Beakers in Bulk.—The first ten solutions are poured into beakers until two-thirds full, and the latter are immersed and brought to the temperature of the bath A. When the first five solutions have been measured, they are taken out of the water-bath and the second series of five beakers inserted in their place, bringing at the same time a third series into the water-bath. The second series are measured and so on. Small gummed labels on the outside prove quite satisfactory for numbering the beakers. It is absolutely necessary to

compare the temperature of the solutions in the beakers with the waterbath from time to time.

After each immersion, the prism should be wiped dry with a clean, soft piece of old linen.

Investigations of Solutions Excluded from Air.—Quickly evaporating liquids, for instance ether solutions, should be treated individually by means of the metal beaker adapted to fit the prism end of the refractometer. To fill the beaker, the refractometer is held in the left hand with the prism pointing upwards, and the metal beaker (M, Fig. 40) is set and securely fastened by means of the bayonet joint. It is now filled quite full and the cover D carefully fitted and locked.

The solution is now enclosed, air and water tight. The refractometer as before is hung upon the wire frame of the bath, with the metal beaker submerged in the bath.

It is expedient to place the solutions before investigation in closed flasks in the nine unoccupied holes in the bath.

After the measurement, the refractometer is held in the left hand with the prism pointing downwards, and the beaker together with its cover detached by giving a slight turn with the right hand. The solution can be used for other purposes, since it has undergone no change in constitution. Finally, the cover is detached from the beaker, and cover, beaker, and prism cleaned by simple means, and the refractometer made ready for the reception of the next solution, as above.

Investigations of Small Quantities of Solutions with the Auxiliary Prism.—When the solution does not occur in sufficiently large quantities for investigation in the glass beaker, or when the solution is too deeply colored, as in dark beers, molasses, etc., the auxiliary prism is used. As described under "Solutions Excluded from Air," the metal beaker without cover is fitted on the refractometer. The auxiliary prism is held horizontally, and, a few drops of the solution having been applied to the hypothenuse face, the prism is inserted into the metal beaker, held conveniently for the purpose, with its hypothenuse face laid upon the polished elliptical face of the refractometer prism, and then locked in by the cover. If an insufficient quantity of the solution has been taken, the margins of the out-spread drops lying between the two prisms can be recognized by looking through the window of the cover on which abuts the square polished end of the auxiliary prism. It is strongly recommended, wherever possible, to apply a sufficiency of the solution, so that the space between the prisms is completely filled, otherwise a loss in brilliancy occurs, and, under certain circumstances, an unavoidable

TABLE OF INDICES OF REFRACTION, n_{D^*} (Compared with Scale Readings of Zeiss Immersion Refractometer, according to Wagner.)

(Com	pared with	ocaic .	readings o	1 72035	IIIIIIICI SIOII	Remaci	onieter, act	ording	to wagner.)
Scale Read- ing.	*D*	Scale Read- ing.	*D.	Scale Read- ing.	* _D .	Scale Read- ing.	* _D .	Scale Read- ing.	n _D .
0.0	1.327360	5.0	1.329320	10.0	1.331260	15.0	1.333200	20.0	1.335168
0.1	1.327399	5.1	1.329350	10.1	1.331299	15.1	1.333238	20.1	1.335168
2	438	2	398	2	388	2	276	2	206
3	477	3	437	3	377	3	314	3	244
4	516	4	476	4	416	4	352	4	282
5 6	555	5 6	515	5 6	455	5 6	390 428	5 6	320
7	594 633		554 593		494 533	2	466		358 396
7 8	672	7 8	632	7 8	572	7 8	504	7 8	434
9	711	9	671	9	611	9	542	9	472
ı.ó	750	6.ó	710	11.ó	650	16.ó	580	21.0	510
1.1	1.327789	6.1	1.329749	11.1	1.331689	16.1	1.333619	21.1	1.335549
2	828	2	788	2	728	2	658	2	588
3	867 906	3		3	767 806	3	697	3	627 666
. 4	945	4	905	4	845	4	736	4	705
5	984	5	944	5	884	5 6	775	5 6	744
	1.328023		982		932	7	833		783
7 8	062	7 8	1.330022	7 8	96 ₂	7 8	892	7 8	822
9	101	7.0	100	9	1.332001	9	931	9	861
2.0	140	7.0	100	12.0	040	17.0	970	22.0	900
2.1	1.328180	7.1	1.330139	12.1	1.332078	17.1	1.334008	22.1	1.335938
2	220	2	178	2	116	2	046	2	976
3	657	3	217	3	154	3	084	3	1.336014
4	300 340	4	256 . 295 .	4	192 230	4	122 160	4	052 000
5 6	380	5	334	5 6	268	5 6	108	5 6	128
7	420	7	373		304		236		166
7 8	460	7 8	412	7 8	344	. 7 8	274	7 8	204
9	500	9	451	9	382	9	312	9	242
3.0	540	8.0	490	13.0	420	18.c	350	23.0	280
3.1	1.328579	8.1	1.330528	13.1	1.332459	18.1	1.334389	23.1	1.336319
2	618	2	566	2	498	2	428	2	358
3	657 696	3	604 642	3 4	537 576	3 4	467 506	3	397 436
+ +	735	7	680		615	7	545	4	475
5	774	5	718	5	654	5 6	584	5 6	514
7 8	813	7 8	756	7 8	693	7 8	623	7 8	553
	852		794	8	732		662	8	592
9	891	9	832	9	771 810	9	701	9	631
4.0	930	9.0	870	14.0	810	19.0	740	24.0	670
4. I 2	1.328969	9. I 2	1.330909 948	14.1	t . 332849 888	19.1	1.334770 818	24.1	1.336708
3	1.329008	3	948 987		927	3	857	2	746 784
4	085		1.331026	; 3	966	3	896	3	822
5	125	4 5 6	104		1.333005		935		860
5 6	164		101	5 6	014	5 6	974	5	898
7 8	203	7 8	143	7 8	083	7 8	1.335013	7 8	936
	242		182	1	122		052		974
9	281	9	221	9	161	9	091	9	1.337012
5.0	320	10.0	260	15.0	200	20.0	130	25.0	050
<u>. </u>	I	1	1	l		li	l		ــــــ

TABLE OF INDICES OF REFRACTION, n_D —(Continued).

	,		,			T			
Scale Read- ing.	** _D -	Scale Read- ing.	*D-	Scale Read- ing.	*D.	Scale Read- ing.	# _D .	Scale Read- ing.	% D.
25.0	1.337050	30.0	1.338960	35.0	1.340860	40.0	1.342750	45.0	1.344630
25.1	1.337088	30.1	1.338998	35.1	1.340898	40.1	1.342788	45.1	1.344667
2	126	2	1.339036	2	9,36	2	826	2	704
3	164	3	074	3	974	3	864	3	741
4	202	4	112	4	1.341012	4	902	4	778
5 6	240	5 6	150 188	5	050	5 6	940	5	818
0	278			6	088 126	0	978		852
7 8	316	7 8	226 264	7 8	164	7 8	1.343016	7 8	889
	354	11		I .	202	1	054		926
9 26.0	392 430	31.0	302 340	36.0	240	41.0	092 130	46.0	963 1.345coo
2 6. 1	1.337468	31.1	1.339378	36.1	1.341278	41.1	1.343167	46.1	1.345037
2	506	2	416	2	316	2	204	2	074
3	544	3	454	3	354	3	241	3	111
4	582	4	492	4	392	4	278	4	148
5	620	5 6	530	5 6	430	5 6	315	5	185
	658		568		468		352		222
7 8	696	7 8	606	7 8	506	7 8	389	7 8	259
	734	u	644 682	1	544		426	1	296
9	772	9		9	582 620	9	463	9	333
27.0	810	32.0	720	37.0		42.0	500	47.0	370
27. I	1.337849	32.1	1.339758	37.1	1.341657	42.I	1.343538	47.1	1.345408
2	888	2	706	2	694	2	576 614	2	446
3	927	3	834	3	731	3	014	3	484
4	966	4	872	4	768	4	652	4	522
5	1.338005	5	910	5 6	805 842	5	690	5	560
	044		948 986		879		728 766		598
7 8	083	7 8		7 8	916	7 8	804	7 8	636 674
9	161	9	1.340024	9	953	9	842	9	712
28.0	200	33.0	100	38.0	990	43.0	880	48.0	750
28. I	τ. 338238	33. т	1.340138	38.1	1.342028	43.1	1.343918	48.1	1.345787
2	276	2	176	2	o66 l	2	956	2	824
3	314	3	214	3	104	3	994	3	861
4	352	4	252	4	142	4	1.344032	4	898
5	390	5 6	290	5	180	5 6	070	5 6	935
0	428		328		218		108		972
7 8	466	7 8	366	7 8	256	7 8	146 184	7 8	1.346009
9	504 542	9	404 442	9	294 332	9	222	9	o83
29 .0	580	34.0	480	39.0	370	44.0	260	49.0	120
29.1	1.338618	34.1	1.340518	39.1	1.342408	44.1	1.344297	49.1	1.346158
2	656	2	556	2	446	2	334	2	106
3	694	3	594	3	484	3	371	3	234
4	732	4	632	4	522	4	408	4	272
5 6	770	5 6	670	5 6	560	5 6	445	5	310
6	808		708		598		482		348
7 8	846	7 8	746	7 8	636	7 8	519	7 8	386
	884	11	784		674	1	556	1	424
9	922	25 0	822 860	40.0	712 750	45.0	593 630	50.0	462 500
30. 0	, ,,,,	35.0	550	40.0	/30	43.0	030	30.0	, , ,
		"						·	·

TABLE OF INDICES OF REFRACTION, n_D —(Continued).

		n		,				·	
Scale Read- ing.	* _D .	Scale Read- ing.	*D*	Scale Read- ing.	- # _D -	Scale Read- ing.	* _D .	Scale Read- ing.	* _D .
50.0	1.346500	55.0	1.348360	60.0	1.350210	65.0	1.352050	70.0	1.353880
50.1	1.346537	55.1	1.348397	60. r	1.350247	65.1	1.352087	70.1	1.353917
2	574 611	2	434	2	284	2	124	2	954
3	611	3	471	3	321	3	161	3	991
4	648	4	508	4	358	4	198	4	1.354028
5 6	685	5	545	5	395	5 6	235	5 6	065
	722	0	582		432	0	272		102
7 8	759	7 8	619	7 8	469	7 8	309	7 8	139
	796	11	656	11	506		346	11	176
9	833 870	56.0	693	61.0	543 580	66.0	383 420	9	213
51.0	1	11	730			i		71.0	250
51.1	1.346907	56.1	1.348767	61.1	1.350617	66.1	1.352457	71.1	1.354286
2	944	2	804	2	654	2	494	2	322
3	981	3	841	3	691	3	531 568	3	358
4	1.347018	4	878	4	728 765	4	506	4	394
5	O55 O92	5 6	915	5	802	5	605 642	5 6	430
-	129		952 989		839		670		466 502
7 8	166	7 8	1.349026	7 8	876	7 8	716	7 8	538
	203	9	063	9	913	9	753	9	574
52.0	240	57.0	100	62.0	950	67.0	790	72.0	610
52.1	1.347277	57.1	1.349137	62.1	1.350987	67.1	1.352827	72.1	1.354646
2	314	2	174	2	1.351024	2	864	2	682
3		3	211	3	061	3	901	3	718
4	351 388	4	248	4	098	4	938	4	754
4 5 6	425	5	285	5 6	135	5	975	5 6	700
ő	462	ő	312	6	172	ő	1.353012		826
7 8	499	7 8	350	7 8	209	7 8	049	7 8	862
8	536	8	396	8	246	8	o 86	∥ 8	898
9	573	9	433	9	283	9	123	9	934
53.0	610	58.ó	470	63.0	320	68.0	160	73.0	970
53.1	1.347647	58.1	1.349507	63. I	1.351357	68.1	1.353196	73.1	1.355006
2	684	2	544	2	394	2	232	2	042
3	721	3	581	3	431	3	268	3	078
4	758	4	618	4	468	4	304	4	114
5	795	5 6	655	5 6	505	5 6	340	5	150
	832		692		542		376	ó	186
7 8	869	8	729	. 7	579 616	7 8	412	7 8	222
	906		766	1			448	11	258
9	943	9	803	9	653	, 9	484	9	294
54.0	980	59.0	840	64.0	690	69.0	520	74.0	330
54.1	1.348018	59.1	1.349877	64.1	1.351726	69.1	1.353556	74.1	1.355366
2	056	2	914	2	762 798	2	592 628	2	402 438
3	094	3	951 988	3 4	798 834	3	664	3 4	430 474
4	132	4			870	4	700		474 510
5	170 208	5	1.350025 062	5 6	906	5 6	736	5	546
	206 246		002		942		730		582
7 8	284	7 8	136	7 8	978	7 8	772 808	7 8	618
	322	9	173	9	1.352014	9	844	9	659
55.0	360	60.ó	210	65.0	050	70.0	880	75-0	690
			J						

TABLE OF INDICES OF REFRACTION, n_D —(Continued).

Scale Read- ing.	₩ _D .	Scale Read- ing.	* _D .	Scale Read- ing.	*D.	Scale Read- ing.	* _D .	Scale Read- ing.	# _D .
75.0	1.355690	80.0	1.357500	85.0	1.359300	90.0	1.361090	95.0	1.362870
75.1	1.355727	80. r	1.357536	85.1	1.359336	90.1	1.361126	95.1	1.362906
2	764	2	572 608	2	372	2	162	2	942
3	801	3	608	3	408	3	198	3	978
4	838	4	644	4	444	4	234	4	1.363014
5 6	875	5 6	68o	5	480	5	270	5 6	050
. 6	912		716	6	516		306		o86
7 8	949	7 8	752	7	552 588	7 8	342	7 8	122
	986	It	788	8	588		378	F1	158
. 9	1.356023	9	824	0, 9	624	9	414	, 9	194
76.0	060	81.0	860	86.0	660	91.0	450	96.0	230
76. ı 2	1.356096	81.1	1.357896	86. r	1.359696	91.1	1.361486	96.1	1.363256
	132 168	14	932 968)	732 768	i	522	2	292
3	204	3	1.358004	3	804	3	558	3 4	328
4	240	4	040	4	840	4	594 630		364 400
· 6	276	5	076	5 6	876	5 6	666	5 6	436
7	312		112		912		702		430
7 8	348	7 8	148	7 8	948	7 8	738	7 8	518
9	384	9	184	9	984	9	774	9	554
77.0	420	82.0	220	87.0	1.360020	92.0	810	97.ó	590
77.1	1.356456	82.1	1.358256	87.1	1.360056	92.1	1 361846	97.I	1.363625
2	492	2	292	2	092	2	882	2	660
3	528	3	328	3	128	3	.918	3	695
4	564	4	364	4	164	4	954	4	730
5 6	600	5 6	400	5	200	5 6	990	5 6	765
	636		436	6	236		1.362026		800
7 8	672	7 8	472	7 8	272	7 8	062	7 8	835
	708	11	508	1	308	1	098	11	870
78.0	744 780	83.0	544 580	88.0	344 380	93.0	134 170	98.0	905 940
78. ı	1.356816	83.1	1.358616	88.1	1.360416	93.1	1.362205	98. I	1.363975
2	852	2	652	2	452	2	240	2	1.364010
3	852 888	3	652 688	3	488	3	275	3	045
4	924	4	724	4	524	4	310	4	080
5	960	5	760	5	560	5 6	345	5 6	115
5 6	996		796		506		380		16ŏ
7 8	1.357032	7 8	832	7 8	632	7 8	415	7 8	195
	068	11	868	ı	668		450	II i	230
9	104	9	904	9	704	9	485	9	265
79.0	140	84.0	940	89.0	740	94.0	520	99.0	290
7 9.1	1.357176	84.1	1.358976	89.1	1.360775	94.1	1 362555	99.1	1.364325
2	212	2	1.359012	2	810	2	590	2	360
3	248	3	048	3	845	3	625	3	395
4	284	4	084	4	88o	4	660	4	430
5	320	5 6	120	5 6	915	5 6	695	5	465
0	356		156	1	950		730		500
7	392	7 8	192 228	7 8	985	7 8	765 800	7 8	535
	428	14	228 264		1.361020	14	835	9	570
9 Bo.o	464 500	85.0	300	90.0	055	95.0	870	100.0	605 640

degradation of the sharpness of the border-line. On the other hand, with a sufficient quantity of solution, the border-line is surprisingly sharp.

The refractometer is now suspended on the frame, and the measurement proceeded with as before described. After measurement, the cover is first removed, and the prism allowed to fall into the hollow of the hand, then the beaker is removed to enable the refractometer to be conveniently cleaned.

Strengths of Various Solutions.—The most extensive work on the quantitative determination of the strength of a large number of common aqueous solutions with the immersion refractometer has been done by Wagner, who has published a large number of tables. These tables show the percentage strength (grams per 100 cc. at 17.5° C.) of a large number of salt solutions and of acids, corresponding to the range of scale readings of the instrument, as well as of cane sugar, dextrose, formalde-

SCALE READINGS ON IMMERSION REFRACTOMETER OF VARIOUS STAND-ARD REAGENTS USED IN VOLUMETRIC ANALYSIS.*

	Temperature C.								
	15°.	16°.	17°.	17.5°.	18°.	19°.	20°.	210.	23°.
Hydrochloric acid:						i			
Normal	37 - 45	37.20	36.95	36.85	36.70	36.45	36.20	35.95	35 - 79
Tenth-normal	17.80	17.60	17.40	17.30	17.20	17.00	16.80	16.55	16.30
Sulphuric acid:	1	1	1			•		1	Ū
Normal	30.60	30.40	30.20	30.10	20.05	20.75	20.50	29.25	20.00
Fifth-normal	18.75	18.60	18.40	18.30	18.20	18.00	17.80	17.55	17.30
Tenth-normal								15.90	
Oxalic acid:	1-7:-3	3	173		33	35		13.3.	-33
Half-normal.	22.45	22.30	22.10	22.00	21.00	21.70	21.50	21.25	21.00
Tenth-normal								15.90	
Potassium bitartrate:	127:23	10.93	120.73	20.03	120.33	1	10.13	13.90	23.03
Tenth-normal	17.75	17.55	17.25	17.25	17.15	16.05	16.75	16.50	16.25
Potassium hydroxide:	1.7.73	1.33	1.33	-7.23	1.7.23	10.93	10.73	10.30	10.23
Normal	42 00	42 65	42 40	42 25	42 70	42 80	42 50	42.20	47 05
Tenth-normal	18 45	18 20	18 10	18 00	17 00	17 70	17 50	17.25	17.00
Sodium hydroxide:	10.45	10.30	10.10	10.00	17.90	17.70	17.30	117.23	17.00
Tenth-normal	18 50	18 25		-8 05	77.0-			17.30	
	10.50	10.35	10.15	10.05	17.11	17.75	17.33	17.30	17.05
Sodium thiosulphate;			00 85	00 00	00.65				
Tenth-normal Potassium bichromate:	24.20	24.05	23.05	23.75	23.05	23.45	23.20	22.95	22.70
	l					-6 05	-6	16.50	-6
Tenth-normal	17.75	17.55	17.35	17.25	17.15	10.95	10.75	10.50	10.25
Silver nitrate:		I			4-	l			-0
Tenth-normal	20.20	20.05	19.05	19.75	19.05	19.45	19.25	19.00	10.75
Sodium chloride:			0-						_4
Tenth-normal	10.20	10.00	17.80	17.70	17.00	17.40	17.20	16.95	10.70
Ammonium sulphocyanate:	1 .		l					l	
Tenth-normal	20.00	20.45	20.25	20.15	120.05	19.85	19.05	19.40	19.15

^{*} According to Wagner, all these solutions were made up at 17.5° C. Readings at different temperatures are given for convenience.

hyde, alcohol, etc. All these observations have been based on the Mohr liter, at a temperature of 17.5°. More convenient for the American analyst would be tables based on the use of a higher temperature, say 20°, and the analyst is recommended to work out his own standards for comparison, at the temperature best suited to his special locality and convenience. The instrument is especially useful in preparing normal and tenth-normal solutions.

The table on page 120, from Wagner, shows the strength of various common laboratory reagents.

SCALE READINGS AT TEMPERATURES FROM 10-30° C. Corrected to 17.5°, According to Wagner.

No.	1.	2.	3.	4.	5.	6.	7.	8.	9.	10.	11.	12 & 13.	No.
,					Scale	Readi	ng at	7.5° C				<u> </u>	-gra-
Tempera- ture C.	15.	20.	25.	30.	35.	40.	45.	50.	60.	70.	80.	90 & 100.	Tempera- ture C.
30	-3.20	3.15	3.25	3.40	3.55	3.65	3.90	4.05	4.20	4.60	4.80	5.25	30
29 28	2.90	2.85	2.95 2.65	3.10 2.80	3.25 2.95	3·35 3·05	3·55 3·25	3·75 3·45	3.90 3.60	4.25 3.90	4·45 4·10	4.50	29 28
27 26	1.00	2.25 1.95	2.35 2.05	2.50	2.65 2.35	2.75 2.45	2.95 2.55	3.15 2.80	3.30 2.95	3.50 3.10	3·75 3·30	4.10 3.65	27 26
25	1.75	1.75	1.80	1.90	2.05	2.15	2.25	2.45	2.60	2.70	2.95	3.20	25
24 23 22 21	1 50 1 25 1,00	1.45 1.25 1.00	1.55 1.30 1.05 0.80	1.60 1.35 1.10 0.85	1.45 1.15	1.85 1.55 1.25 0.95	1.95 1.65 1.30	2.10 1.75 1.40 1.05	2.25 1.90 1.55 1.20	2.35 2.00 1.65 1.25	2.55 2.15 1.75 1.35	2.75 2.35 1.90 1.45	24 23 22 21
20	0.50	0.50	0.55	0.60	0.65	0.65	0.75	0.75	0.85	0.90	0.95	1.05	20
19	0.30	_		0.35 0.15	0.40	0.40		0.45	0.45	0.55	0.55	0.60	19
17.5	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	17.5
17 16	-0.10 0.30					0.10	0.15			0.15 0.50		ı	17 16
15	0.50	0.45	0.45	0.50	0.60	0.60	0.65	0.75	0.75	0.80	0.85	0.90	15
14 13 12 11	0.70 0.85 1.00 1.15					0.85		, ,					14 13 12 11
10	1.25												
No.	1.	2.	3.	4.	5.	6.	7.	8.	9.	10.	II.	:2 & 13.	No.

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CHAPTER VII.

MILK AND ITS PRODUCTS.

MILK.

Nature and Composition.—Milk is the secretion of the mammary glands of female mammals for the nourishment of their young. Containing as it does all the requisites for a complete food, i.e., sugar, fat, proteins, and mineral ingredients, combined in appropriate proportion, there is ample reason why it occupies so high a place in the scale of human foods. It is a yellowish-white opaque fluid, denser than water, containing in complete solution the sugar, soluble albumin, and mineral content, and, in less complete solution, the casein, while the fat-globules are held in suspension in the serum, forming an emulsion.

The specific gravity of pure milk ranges from 1.027 to 1.035.

Milk from various animals has the same general physical properties and the same ingredients, differing, however, in percentage composition. Of all the varieties, the milk of the cow is by far the most important from its universal use, and, unless otherwise qualified, the term milk wherever it occurs in this volume will be understood to mean cow's milk.

Acidity.—When perfectly fresh, milk of carnivorous mammals is, as a rule, acid in reaction, while human milk and that of the herbivora is alkaline. Cow's milk, when freshly drawn, is more often amphoteric in reaction, i.e., it reacts acid with blue and alkaline with red litmus. It soon becomes distinctly acid, and the acidity increases as the milk sugar gradually becomes converted into lactic acid.

Microscopical Appearance.—Under the microscope pure milk shows a conglomeration of various-sized fat globules having a pearly lustre. These globules vary from 0.001 to 0.01 mm. in diameter, averaging about 0.005 mm. When examined under very high powers, it is possible to distinguish bacteria in the milk, the number to be seen depending greatly on the time that has elapsed since the milk was drawn from its source, as well as on the surroundings, the conditions of handling, exposure, etc.

Color.—The yellow color of milk is imparted to it by the fat globules, and varies greatly in milk from different breeds of cattle, as well as in milk from the same cow at different seasons, being, as a rule, paler during the winter or stall-fed months, and having its greatest intensity soon after the cow is put out to pasture.

Milk Sugar, the carbohydrate of milk, is normally present in amounts varying from 3 to 5 per cent. For the properties of milk sugar see page 577.

The Proteins of Milk.—Casein constitutes about 80% of the entire proteins of milk, being present in an average sample to the extent of about 3%. It exists in combination with calcium phosphate, and probably does not form a perfect solution in the milk, but is rather diffused therein in a somewhat colloidal form, being so finely divided, however, as to be incapable of separation by filtration while the milk is fresh.

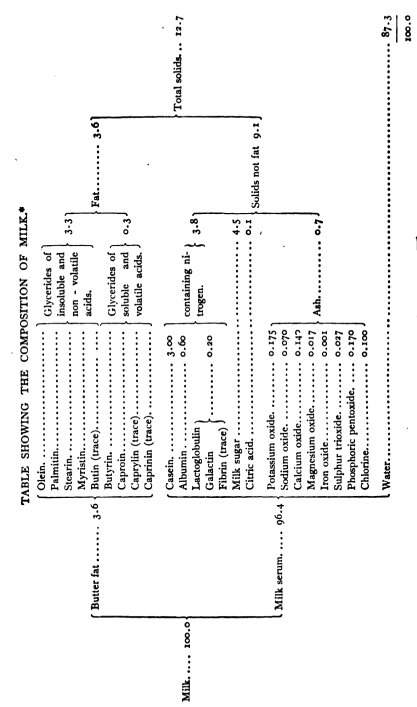
Pure casein is a white, odorless, and tasteless solid, sparingly soluble in water, and insoluble in ether and alcohol. It is readily soluble in dilute alkalies. Strong acids also dissolve it, but its character is changed. From alkaline solution it is precipitated without change by neutralizing with acid. Its solutions are lævo-rotary.

Lact-albumin is the soluble albumin of milk, existing therein to the extent of about 0.6% and forming about 15% or more of the milk proteins. It much resembles the albumin of eggs, being coagulated at 70° to 75° C. It is readily soluble in water. Its specific rotary power according to Béchamp is $[\alpha]_D = -67.5$.

Lactoglobulin has been discovered by Emmerling as a constituent in milk, but exists in traces only. According to Babcock, it may be separated from milk whey by carefully neutralizing with sodium hydroxide, and afterwards saturating with magnesium sulphate. It much resembles the globulin of blood serum, being coagulated at 67° to 76° C.

Fibrin.—Babcock has discovered in milk very minute traces of a substance analogous to the fibrin of blood. This substance, it is claimed, forms a part of the slime found in the separator-bowl of a centrifugal skimmer.

Other Nitrogenous Substances.—Besides the above normal constituents of milk. certain bodies may be formed by proteolytic action during fermentation, such, for example, as caseoses and peptones, formed for the most part by the decomposition of a part of the casein. Galactin is a gelatin-like body of the nature of peptone, occurring in traces in milk. Besides these, minute traces of amido-bodies, such as creatin and urea, are sometimes present, and also ammonia.



* According to S. M. Babcock.

Milk Fat.—Fat forms the most variable constituent of milk, being found in proportions ranging from 2.5 to 7 per cent. For the chemical composition and characteristics of milk fats see Butter (p. 529).

The fat globules are held in suspension in the milk and have long been thought to be surrounded each by a thin nitrogenous membrane, known as *Storch's mucoid protein*, which becomes broken on churning. This theory, while rendered probable by many of the phenomena connected with the dairy, is by no means universally held at present.

Citric Acid has been found to exist in milk, probably in combination with certain of the mineral constituents, being present to the extent of about 0.1%.

The table on page 126 arranged by Babcock shows quite clearly the percentage composition of an average cow's milk.

For comparison of milk from different animals the following table * is inserted, showing in most cases minimum, maximum, and mean determinations from a large number of actual analyses:

	No. of Anal- yses.	Specific Gravity.	Water.	Casein.	Albu- min.	Total Pro- teids.	Fat.	Milk Sugar.	Ash.
Cow's milk	800								
Minimum		1.0264	80.32	1.79	0.25	2.07	1.67	2.11	0.35
Maximum		1.0370	90.32	6.29	1.44	6.40	6.47	6.12	1.21
Mean		1.0315	87.27	3.02	0.53	3-55	3.64	4.88	0.71
Human milk	200		' '						
Minimum	1	1.027	81.09	0.18	0.32	0.69	1.43	3.88	0.12
Maximum	1	1.032	91.40	1.96	2.36	4.70	6.83	8.34	1.90
Mean		-	87.41	1.03	1.26	2.29	3.78	6.21	0.31
Goat's milk	200		•		i			1	
Minimum		1.0280	82.02	2.44	0.78		3.10	3.26	0.39
Maximum		1.0360	90.16	3.94	2.01		7-55	5-77	1.06
Mean		1.0305	85.71	3.20	1.09	4.29	4.78	4.46	0.76
Ewe's milk	32				-			1	-
Minimum		1.0298	74.47	3.59	0.83	. .	2.81	2.76	0.13
Maximum		1.0385	87.02	5.69	1.77	[]	9.80	7.95	1.72
Mean		1.0341	80.82	4.97	1.55	6.52	6.86	4.91	0.89
Mare's milk	47								-
Mean		1.0347	90.78	1.24	0.75	1.99	I.2I	5.67	0.35
Ass's milk	5					''			•
Mean		1.036	89.64	0.67	1.55	2.22	1.64	5-99	0.51

Composition of the Ash of Milk.—The ash of milk does not truly represent the mineral content, since, in the process of incineration, the character of some of the constituents is altered by oxidation and otherwise.

Expressed in parts per 100, the ash of the typical milk sample whose full analysis is given on page 126 would be about as follows:

^{*} Compiled from König's Chemie der mens. Nahr. u. Genuss.

Potassium o	xide		 • • • •	 		25.02
Sodium	"		 ••••	 • • • •	• • • • •	10.01
Calcium	"		 ••••	 		20.01
Magnesium	"		 	 		2.42
Iron	"		 	 		0.13
Sulphur trio	xide	• • • • •	 	 		3.84
Phosphoric 1	pent	oxide.	 	 	• • • • •	24.29
Chlorine	- 		 	 	• • • • •	14.28
						100.00

Soldner regards the following as more nearly representing the proportion in which the mineral salts exist in milk:

	Per Cent.
Sodium chloride, NaCl	10.62
Potassium chloride, KCl	9.16
Mono-potassium phosphate, KH ₂ PO ₄	12.77
Di-potassium phosphate, K ₂ HPO ₄	9.22
Potassium citrate, $K_3(C_6H_5O_7)_2$	5-47
Di-magnesium phosphate, MgHPO	3.71
Magnesium citrate, Mg ₃ (C ₆ H ₅ O ₇) ₂	4.05
Di-calcium phosphate, CaHPO	7-42
Tri-calcium phosphate, Ca ₃ (PO ₄) ₂	8.90
Calcium citrate, Ca ₃ (C ₆ H ₅ O ₇) ₂	23-55
Lime, combined with proteins	5.13
	100.00

Fore Milk and Strippings.—Unless a portion drawn from the well-mixed or whole complete milking of an animal is taken for analysis, one does not get a fair representative sample of the milk, for it is a well-known fact that the first portion of milk drawn from the udder, termed the "fore milk," is very low in fat, while the last portions or "strippings" contain a very high fat content, sometimes exceeding 10% fat. The following analyses show the difference between fore milk and strippings in two cases:

	Per Cent Water.	Per Cent Solids.	Per Cent Fat.
(1) Fore milk. Strippings. (2) Fore milk. Strippings.	80.82 88.73	11.83 19.18 11.27 19.63	1.32 9.63 1.07 10.36
			!

The per cent of albuminoids, sugar, and ash is nearly the same in both fore milk and strippings.

Colostrum.—The milk given by cows and other mammals for two or three days after the birth of young is termed colostrum, and differs materially in composition from normal milk. It is yellow in color, of an oily consistency, and has a pungent taste. It acts as a purge upon the young. Examined under the microscope, it is found to contain large circular cells larger than fat globules and somewhat similar to blood corpuscles. It is very high in albumin, which seems to be similar to blood albumin. The following analyses were made by Engling, showing the composition of colostrum from a cow eight years old:

Time after Calving.	Specific Gravity.	Fat.	Casein.	Albu- min.	Sugar.	Ash.	Total Solids.
Immediately. After 10 hours. 24 48 72 48 72 48 48 48 48 48 48 48 48 48 4	1.068 1.046 1.043 1.042 1.035	3-54 4.66 4-75 4.21 4.08	2.65 4.28 4.50 3.25 3.33	16.56 9.32 6.25 2.31 1.03	3.00 1.42 2.85 3.46 4.10	1.18 1.55 1.02 0.96 0.82	26.93 21.23 19.37 14.19

The average of twenty-two analyses of colostrum from different cows by Engling showed total solids 28.31, fat 3.37, casein 4.83, albumin 15.85, sugar 2.48, ash 1.78.

Frozen Milk.—Since it is the water in milk that freezes, it follows that in partially frozen milk the unfrozen portion of the milk, or that part which remains still liquid, becomes concentrated by the process of freezing. This is borne out by the following figures of Richmond:*

	Frozen Portion, Per Cent.	Unfrozen Portion, Per Cent.
Water	96.23	85.62
Fat	1.23	4.73
Sugar	I.42	4.95
Proteins	91	3.90
Ash	21	.80
Specific gravity	1.0090	1.0345

Fermentations of Milk.—These are due to the action of bacteria of various kinds, the most common being the lactic fermentation.

The Souring of Milk is caused by the action of a large number of species of acid-forming bacteria, chief among which is the Bacillus acidi lactici, which multiplies faster than other bacteria in raw milk under

^{*} Analyst, XVIII. p. 53.

favorable conditions of temperature. Part of the milk sugar is acted on and transformed, first into dextrose and galactose, the latter sugar subsequently forming lactic acid, as follows:

(1)
$$C_{12}H_{22}O_{11},H_2O = C_6H_{12}O_6 + C_6H_{12}O_6$$
Lactose Dextrose Galactose

(2)
$$C_6H_{12}O_6 = 2C_3H_6O_8$$
Galactose Lactic acid

More and more acid is formed until the casein can no longer be held up, curdling ensues, and the casein is precipitated. Finally, after a certain degree of acidity is reached, the ferment is killed and the action stops. Other acids than lactic are also undoubtedly produced, since a small part of the acid in sour milk is found to be volatile. According to Conn * the volatile acids are acetic and formic.

Abnormal Fermentation.—Through the agency of micro-organisms that may develop under certain conditions, various changes are produced in milk that to some extent alter its character. Thus bitter milk is sometimes produced as the result of some organism as yet but little understood.

Occasionally milk is found possessing a peculiarly thick and slimy consistency, whereby it may be drawn out in threads, by dipping a spoon into the milk and withdrawing it therefrom. This is termed ropy milk, and is more often met with in warm weather. It is undoubtedly produced as a result of bacterial action.

Enzyme-forming Bacteria are not uncommonly developed in milk, causing various proteolytic changes, whereby the casein is partially transformed into peptones, caseoses, etc.

Chromogenic Bacteria are the agencies that produce peculiar pigments in milk. Thus red milk is due to Bacillus erythrogenes; yellow milk to Bacillus xynxanthus; blue milk to Bacillus cyanagenes. The latter is quite common, appearing ordinarily in patches in the milk.

CHEMICAL ANALYSIS OF MILK.

Ordinarily, in ascertaining the nutritive value of milk, one determines its specific gravity, total solids, fat, protein, milk sugar, and ash. Occasionally it is thought desirable to make a distinction in the case of protein between the casein and the albumin. Rarely is it necessary to further subdivide the nitrogenous bodies in milk, unless in connection with a special study of the proteolytic changes which it undergoes.

The total solids, fat, and ash are usually all determined directly, and,

^{*} U. S. Dept. of Agric., Off. of Exp. Stations, Bul. 25, p. 21.

in the case of the milk sugar and the proteins, a determination of either one may be directly made (whichever is most convenient), the other being calculated by difference.

When foreign ingredients or adulterants are present in milk, special methods are employed to detect them.

Preparation of the Sample.—In procuring a sample for analysis, the greatest care is necessary to insure a homogeneous sample. By far the best method in every case, where possible, is to pour the milk back and forth from one vessel to another (i.e., pour from the original container into an empty vessel and back at least once). Where this is impossible from the size of the container or for any other reason, the milk should be thoroughly mixed with a dipper. A "sampler," of which the Scovell sampling-tube (Fig. 41) is a convenient form, also aids in securing a representative sample, and is invaluable when it is desirable to secure a definite fraction of the whole for a composite sample.

This instrument consists of a brass or copper tube made in two parts which telescope accurately together as shown in Fig. 41, the lower part being closed at the bottom, but provided with three or more lateral slits. The sampler, drawn out to its full length, is carefully inserted in the tank containing the milk and lowered to the bottom, after which the upper part is pressed down over the lower so as to close the slits, and the tube is then lifted out of the tank, containing a fairly representative sample of the milk.

In all operations to which a milk sample is submitted during the process of analysis, it should invariably be poured into a clean empty vessel and back, whenever it has been at rest for an appreciable time, in order to insure a homogeneous mixture.

readily obtained with the aid of a hydrometer, accurately graduated within the limits of the widest possible variation in the specific gravity of milk. Hydrometers for vell Milk-sampling special use with milk are known as lactometers, and are graduated variously. One of the most convenient forms Fig. 42.—The Quevenne lactometer, graduated from 15° to 40°, corresponding to specific gravity 1.015 to 1.040. This

instrument, shown in Fig. 42, has a thermometer combined with it, the stem containing a double scale, on the lower part of which the specific gravity is read, while the temperature is read from the upper part.

Another form of instrument is termed the New York Board of Health lactometer, which is not graduated to read the specific gravity directly, but has an arbitrary scale divided into 120 equal parts, the zero being equal to the specific gravity of water, while 100 corresponds to a specific gravity of 1.029. To convert readings on the New York Board of Health scale to Quevenne degrees they must be multiplied by .29.

QUEVENNE LACTOMETER DEGREES CORRESPONDING TO NEW YORK BOARD OF HEALTH LACTOMETER DEGREES.

Board of Health Degrees.	Quevenne Scale.	Board of Health Degrees.	Quevenne Scale.	Board of Health Degrees.	Quevenne Scale.
60 61 62 63 64 65 66 67 68 69 70	17-4 17-7 18-0 18-3 18-6 18-8 19-1 19-4 19-7 20-0 20-3 20-6	81 82 83 84 85 86 87 88 89 90	23.5 23.8 24.1 24.4 24.6 24.9 25.2 25.5 25.8 26.1 26.4	101 102 103 104 105 106 107 108 109 110	29.3 29.6 29.9 30.2 30.5 30.7 31.0 31.3 31.6 31.9 32.2
72 73 74 75 76 77 78 79	20.9 21.2 21.5 21.7 22.0 22.3 22.6 22.9 23.2	93 94 95 96 97 98 99	27.0 27.3 27.6 27.8 28.1 28.4 28.7 29.0	113 114 115 116 117 118 119	32.8 33.1 33.4 33.6 33.9 34.2 34.5 34.8

If extreme accuracy is desired, the Westphal balance or the pycnometer should be used for the determination of specific gravity. For ordinary cases, however, the lactometer, if carefully made, is sufficiently accurate.

With any other form of lactometer than the Quevenne, a separate thermometer is necessary in order to determine the temperature, the common practice being to standardize all such instruments at 60° F. (15.6° C.).

Readings at temperatures other than 60° may be corrected to that temperature by the aid of the table on page 133.

DETERMINATION OF TOTAL SOLIDS.—Dish Method.—For purposes of milk analysis, platinum dishes are by far the most desirable. These, if made for the purpose, should be of the shape shown in Fig. 52, measur-

FOR CORRECTING THE SPECIFIC GRAVITY OF MILK ACCORDING TO TEMPERATURE (BY DR. PAUL VIETH).

Degrees of					D	egrees	of Th	nermo	meter	(Fah	renhei	t).				
Lactom- eter.	45	46	47	48	49	50	51	52	53	54	5.5	56	57	58	59	60
20																
				20.1												
22				21.1												
23																
				23.1												
25	23.8	23.9	24.0	24.0	24.1	24.1	24.2	24-3	24.4	24.5	24.0	24.0	24-7	24.8	24.9	_
2Ğ	24.8	24.9	24.9	25.0	25.1	25.1	25.2	25.2	25.3	25.4	25.5	25.6	25.7	25.8	25.9	-
27	25.8	25.9	25.9	20.0	20. I	20.1	20.2	20.2	20.3	20.4	20.5	20.0	20.7	20.8	26.9	—
	20.7	20.8	20.8	20.9	27.0	27.0	27.1	27.2	27.3	27.4	27.5	27.0	27.7	27.8	27.9	
9	27.7	27.8	27.8	27.9	28.0	28.0	28. I	28.2	28.3	28.4	28.5	28.0	28.7	28.8	28.9	
30	28.0	28.7	28.7	28.8	28.9	29.0	29.1	29.1	29.2	29.3	29.4	29.0	29.7	29.8	29.9	_
3r	29.5	29.0	29.0	29.7	29.8	29.9	30.0	30.1	30.2	30.3	30.4	30.5	30.0	30.8	30.9	_
32	30.4	30.5	30.5	30.0	30-7	30.9	31.0	31.1	31.2	31.3	31.4	31.5	31.0	31.7	31.9	_
3	31.3	31-4	31.4	31.5	31.0	31.8	31.9	32.0	32.1	32-3	32.4	32.5	32.0	32-7	32.9	_
4	32.2	32.3	32.3	34 - 4	32.5	32.7	32.9	33.0	33.1	33-2	33-3	33-5	33.0	33-7	33-9	_
35	33.0	33.1	33.2	33-4	33-5	33.0	33.8	33-9	34.0	34.2	34 - 3	34 - 5	34.0	34 - 7	34-9	_
				-				<u> </u>						!		
		61	62	63	64	65	66	67	68	69	70	71	72	73	74	75
20		20.1	20.2	20.2	20.3	20.4	20.5	20.6	20.7	20.0	21.0	21.1	21.2	21.3	21.5	21.6
21																
22		22.I	22.2	22.3	22.4	22.5	22.6	22.7	22.8	23.0	23.1	23.2	23.3	23.4	23.5	23.7
3		23.I	23.2	23.3	23.4	23.5	23.6	23.7	23.8	24.0	24. I	24.2	24.3	24.4	24.6	24.7
14				24.3												
5		25.I	25.2	25.3	25.4	25.5	25.6	25.7	25.0	2Ď.O	2Ď. I	26.2	26.4	26.5	26.6	26.8
		26. I	26.2	26.3	26.5	26.6	26.7	26.8	27.0	27.I	27.2	27.3	27.4	27.5	27.7	27.8
7]	27.I	27.3	27.4	27.5	27.6	27.7	27.8	28.0	28. I [†]	28.2	28.3	28.4	28.6	28.7	28.0
8]:	28. I	28.3	28.4	28.5	28.6	28.7	28.8	20.0	20. I	20.2	20.4	20.5	20.7	20.8	20.0
9		20.1	20.3	20.4	20.5	20.6	20.8	20.0	30. I	30.2	20. 2	30.4	20.5	20.7	20.0	21.0
30		30.1	30.3	30.4	30.5	30.7	30.8	30.9	31.1	31.2	31.3	31.5	31.6	31.8	31.0	32.1
, , ,		31.2	31.3	31.4	31.5	31.7	31.7	31.8	32.0	32.2	32.4	32.5	32.6	32.8	33.0	33.1
32		32.2	32.3	32.5	32.6	32 - 7	32.9	33.0	33.2	33-3	33.4	33.6	33-7	33.9	34.0	34.2
		33.2	33.3	33.5	33.6	33.8	33.0	34.0	34.2	34.3	34.5	34.6	34.7	34.0	35.1	35.2
4		34.2	34 - 3	34.5	34.6	34.8	34.0	35.0	35.2	35.3	35.5	35.6	35.8	36.0	36. I	36.1
5		35.2	35.3	35-5	35.6	35.8	35.9	36. I	36.2	36.4	36.5	36.7	36.8	37.0	37.2	37.7
	ľ				, J		اردور	1		را د			اردين	٠,٠٠١,	,,	J/ - J

ing about $2\frac{3}{4}$ inches in diameter at the top, and $2\frac{1}{4}$ inches in diameter at the bottom, having carefully rounded rather than square edges, and being $\frac{1}{2}$ inch deep. The bottom is not perfectly flat, but slightly crowned outward. Such a dish will hold about 35 cc.

For purposes of economy it is best to have these dishes spun out with a thick bottom, but with thin sides, not so thin, however, as to be too readily bent.

If platinum dishes cannot be afforded, dishes of porcelain, glass, aluminum, nickel, or even tin may be used, but in all cases should be as thin as practicable.

About 5 cc. of the thoroughly mixed sample of milk are carefully transferred by means of a pipette to a tared dish on the scale-pan, and its

weight accurately determined. The dish with its contents is then transferred to a water-bath, being placed over an opening preferably but little smaller than the diameter of the bottom of the dish, so that as large a surface as possible is in contact with the live steam of the bath. Here it is kept for at least two hours, after which the dish is wiped dry while still hot, transferred to a desiccator, cooled, and weighed.*

Babcock Asbestos Method.†—Provide a hollow cylinder of perforated sheet metal, 60 mm. long and 20 mm. in diameter, closed 5 mm. from one end by a disk of the same material. The perforations should be about 0.7 mm. in diameter and about 0.7 mm. apart. Fill loosely with from 1.5 to 2.5 grams of freshly ignited, woolly asbestos, free from fine and brittle material, cool in a desiccator, and weigh. Introduce a weighed quantity of milk (between 3 and 5 grams), and dry in a wateroven to constant weight, which is usually reached after four hours' heating.

DETERMINATION OF ASH.—The platinum dish containing the milk residue, obtained in the determination of total solids by the dish method described above, is next placed upon a suitable support above a Bunsen flame (a platinum triangle or a ring stand is convenient for this), and the residue is ignited at a dull-red heat to a perfectly white ash, after which it is cooled and weighed.

DETERMINATION OF FAT.—The Adams Method.—Without doubt the most accurate method of fat determination is by extraction with ether. For this purpose a strip of fat-free filter-paper about 2½ inches wide and 22 inches long is rolled into a coil and held in place by a wire as shown in Fig. 43.

Schleicher and Schüll furnish fat-free strips especially for this work, but it is very easy to prepare the strips and extract them with the Soxhlet apparatus.

About 5 grams of milk are run into a beaker with a pipette, and the weight of the beaker and milk are determined. The coil is then introduced into the beaker, holding it by the wire in such a manner that as

^{*} It is a common practice to transfer the milk residue, after a preliminary drying on the water-bath, to an air-oven, kept at a temperature of from 200°, where it is dried to a constant weight; but after an experience in analyzing over 30,000 samples of milk, the author is prepared to state that in his opinion the results obtained by the above method of procedure, using the water-bath alone, are more satisfactory. It is impossible to keep a milk residue at a temperature above 100° for any length of time without its undergoing decomposition, especially as to its sugar content, as is shown by the darkening in color. A milk residue should be nearly pure white, a brownish color showing incipient decomposition. Hence, by continued heating, especially at the temperature of 105°, the residue would continue to lose weight almost indefinitely. If it is thought best to give a final drying in the air-oven, the time should be short and the temperature employed should not in any case exceed 100°.

[†] A. O. A. C. method, U. S. Dept. of Agric., Bur. of Chem., Bul. 107 (rev.), p. 117.

much as possible of the milk is absorbed by the paper. It is often possible to take up almost the last drop of the milk. By then weighing the beaker, the amount of milk absorbed by the coil is determined by difference, and the paper coil is hung up and dried, first in the air and then in the oven, at a temperature not exceeding 100°. Another method of charging the paper coil consists in suspending it by the wire and gradually delivering upon it 5 cc. of the milk from a pipette, the density of the milk being known.

The coil containing the dried residue is then transferred to the Soxhlet extraction apparatus (see p. 63) and subjected to continuous extraction with anhydrous ether for at least two hours, the receiving-flask being first accurately weighed. The tared flask with its contents is freed from all remaining ether, first on the water-bath and finally in the air-oven. It is then cooled and weighed, the increase in weight representing the fat in the amount of milk absorbed by the coil. If there is any doubt about all the fat having been extracted at first, the process of extraction may be continued till there is no longer a gain in weight of the flask. Experience soon shows the length of time necessary for the complete extraction, which of course depends on the degree of heat employed, and the frequency with which the extracting-tube overflows. Two hours is ample

Adams Milkfat Coil.

for most cases, in which the conditions are such that the ether siphons

over from the extraction-tube ten times per hour.

Babcock Asbestos Method.*—Extract the residue from the determination of water by the Babcock asbestos method with anhydrous ether in a continuous extraction apparatus, until all the fat is removed, which usually requires two hours. Evaporate the ether, dry the fat in the extraction flask at the temperature of boiling water, and weigh. The fat may also be determined by difference, drying the extracted cylinders at the temperature of boiling water.

FAT METHODS BASED ON CENTRIFUGAL SEPARATION. — These methods are the most practicable for commercial work and for use by the public analyst, since they are much more rapid, and, if carefully carried out, practically as accurate as the Adams method. They all depend upon the use of a centrifugal machine, having hinged pockets in which are carried graduated bottles, into each of which a measured quantity of milk is introduced. The milk is then subjected to the action of a suitable reagent, which dissolves the casein and liberates the fat in

^{*} A. O. A. C. method, U. S. Dept. of Agric., Bur. of Chem., Bul. 107 (rev.), p. 119.

a pure state, after which, by whirling at a high speed, the pockets are thrown out horizontally and the milk fat driven into the neck of each bottle, where the amount is directly read.

Various processes of this kind, each having its own special adherents, are in extensive use, among which the best known are the Babcock, the Leffman and Beam, the Gerber, and the Stokes.

A résumé of these processes, showing the reagents employed and other comparative data, is thus tabulated by Allen.*

	Babcock.	Leffman- Beam.	Gerber.	Stokes.
Milk. Sulphuric acid, volume. " " specific gravity. Hydrochloric acid. Amyl alcohol.	17.5 cc. 1.831 to 1.834 None	15 cc. 9 cc. 1.85 1.5 cc. 1.5 cc.	1.82 to 1.825 None	15 cc. 13½ cc. 1.82 to 1.83 None 1.5 cc.

The Babcock Process, devised originally for the use of creameries and dairymen, is now extensively employed for fat determination in the laboratory.

It has stood the test of over ten years' successful use in the writer's hands. During this time on various occasions results as determined have been compared with those obtained by the Adams process, and the agreement has been as close as could be expected. The following figures show the results of such comparative determinations made in duplicate on three samples of milk, viz., a whole pure milk, (1) and (2); a watered milk, (3) and (4); and a milk centrifugally skimmed, (5) and (6).

COMPARATIVE FAT DETERMINATION BY ADAMS-SOXHLET AND BY BABCOCK PROCESSES.

		Per Cent of Fat by the Adams-Soxh- let Process.	Per Cent of Fat by the Babcock Process.
A whole milk	(1)	4.27	4.30
A watered milk	(2)	4.28	4-35
	(4)	2.70 2.74	2.70 2.80
A skimmed milk	(5)	0.16	0.15
	(6)	0.14	0.15

The Centrijuge.—Various styles of centrifuge are in use for this process, some driven by hand, some by steam-power, and some by the electric motor, carrying from 4 to 40 bottles. Fig. 44 shows an 8-bottle hand machine, driven by friction gearing, as well as the steam-driven centrifuge in common use in dairies and creameries, which is a 20-bottle machine

^{*} Commercial Organic Analysis, IV. p. 150.

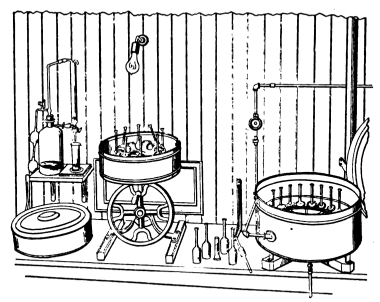


Fig. 44.—Types of the Babcock Centrifuge and Appurtenances. Hand machine at the left; steam-driven machine at the right.

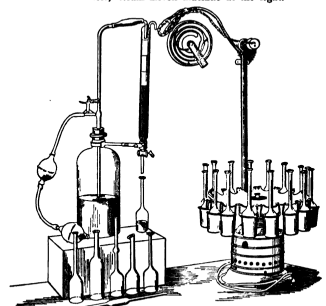


Fig. 45.—Electrically-driven Babcock Centrifuge, with Aluminum Frame, Carrying 16

Bottles. Acid burette at the left.

A sheet-metal safety-shield (removed for showing the construction) normally surrounds the instrument. Such a shield is shown in Fig. 11.

having paddles on the outer periphery of the revolving frame, against which the steam impinges, driving it like a horizontal water-wheel.

The most noiseless and easy-running machine is that driven by an electric motor. An example of this kind of centrifuge is shown in Fig. 45, carrying 16 bottles.

The ordinary Babcock test bottle is shown in Fig. 46, A, that used for skimmed milk in Fig. 46, B. The bottles are graduated with reference to using 18 grams of the sample.

Manipulation.—By means of a pipette graduated to hold 17.6 cc. (the average volume of 18 grams) that amount of the thoroughly mixed sample of milk to be tested is transferred to a test bottle, and 17.5 cc. of commercial sulphuric acid of a specific gravity of 1.82 to 1.84 are added

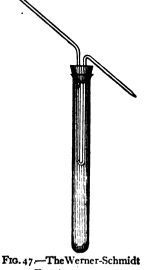


Fig. 46.—Babcock Test Bottles. A, Milk Bottle. B, Skimmed Milk Bottle.

by means of a graduate or an automatic burette, shown in Fig. 44. The contents are then thoroughly mixed, during which operation much heat is developed by the action of the acid on the proteins and milk sugar, and the mixture turns a very dark brown. The test bottles are then placed in the centrifuge pockets (an even number being always used, arranged opposite each other to properly balance) and whirled for at least five minutes. Hot water is then added up to the necks of the bottles, which are then again whirled for about two minutes. Enough hot water is then added to drive the fat into the neck of each bottle, and a final whirl of about a minute's duration is given, after which the bottles are removed from the pockets, and the percentage of fat is read, while still hot, from the graduated neck by means of a pair of calipers.

The Werner-Schmidt Method.—Ten cc. of milk are introduced by means of a pipette into a large test-tube of 50 cc. capacity, and 10 cc. of concentrated hydrochloric acid are added. The mixture is shaken and heated till the liquid turns a dark brown, either by direct boiling for a minute or two, or by immersing the tube in boiling water for from five to

ten minutes. The tube is then cooled by immersion in cold water, and 30 cc. of washed ether is added. The tube is closed by a cork provided with tubes similar to a wash-bottle, the larger tube being adapted to slide up and down in the cork, and preferably being turned up slightly at the bottom. The contents of the tube are shaken, the ether layer allowed to separate, and the sliding-tube arranged so that it terminates slightly above the junction of the two layers. The ether is then blown out into a weighed flask. A second and a third portion of ether of 10 cc. each are successively shaken with the acid liquid and added to the contents of the weighed flask, from which the ether is subsequently evaporated and the weight of the fat easily obtained.



Fat Apparatus.

Instead of measuring the milk into the testingtube, a known weight of milk may be operated on. A sour milk may be readily tested in this way, provided it is previously well mixed.

Determination of Fat by the Wollny Milk-fat Refractometer.*—This instrument presents the same appearance as the butyro-refractometer, Fig. 36, with an arbitrary scale reading from 0 to 100, the equivalent readings in indices of refraction of the Wollny instrument varying from 1.3332 to 1.4220. Exactly 30 cc. of the milk to be tested are measured into the stoppered flask A, Fig. 48. This may be done by the use of the automatic pipette, which holds exactly 7½ cc., removing four pipettes full of the milk. B is a numbered tin sampling-tube in which the milk sample is kept for convenience, and into which the automatic pipette readily fits. Having measured 30 cc. of the milk into the flask A, 12 drops of a solution of 70 grams potassium bichromate and 312.5 cc. of stronger ammonia in one liter of water may be added as a preservative,

^{*} Milch Zeit., 1900, pp. 50-53.

if the sample is to be kept for some time before finishing the test. Twelve drops of glacial acetic acid are added to curdle the milk. The flask is then corked and shaken for one to two minutes in a mechanical shaker, after which 3 cc. of a standard alkaline solution are added, and the flask corked and shaken for ten minutes in the mechanical shaker, the temperature being kept at 17.5° C. The standard alkaline solution is prepared

Fig. 48.- Accessories for Carrying Out the Wollny Milk-fat Process.

by dissolving 800 cc. of potassium hydroxide in a liter of water, adding 600 cc. of glycerin and 200 grams pulverized copper hydrate, the mixture being allowed to stand for several days before using, shaking at intervals. Finally 6 cc. of water-saturated ether are added to the mixture in the flask, using for convenience the automatic pipette fitted in the corked bottle as shown. The flask is again shaken for fifteen minutes in the mechanical shaker, and whirled for three minutes in the centrifuge, after which a few drops of the ether solution are transferred to the refractometer, and the reading taken. The percentage of fat is obtained by means of the following table:

PERCENTAGES OF FAT CORRESPONDING TO SCALE READINGS ON THE WOLLNY REFRACTOMETER.

Scale Read- ing.	Per Cent Fat.	Scale Read- ing.	Per Cent Fat.	Scale Read- ing.	Per Cent Fat.	Scale Read- ing.	Per Cent Fat.	Scale Read- ing.	Per Cent Fat.	Scale Read- ing.	Per Cent Fat.
20.0		24-5	0.41	29.0	0.87	33-5	I - 34	38.0	1.85	42.5	2.41
1 2 3 4		6 7 8 9	0.42 0.43 0.44 0.45	1 2 3 4	0.88 0.89 0.90 0.91	6 7 8 9	1.35 1.36 1.37 1.38	1 2 3 4	1.87 1.88 1.89	6 7 8 9	2.43 2.44 2.46 2.47
5		25.0	0.46	5	0.92	34.0	1.39	5	1.91	43.0	2.49
6 7 8 9 21.0	0.00 0.01 0.02 0.03 0.04	1 2 3 4 5	0.47 0.48 0.49 0.50 0.51	6 7 8 9 30.0	0.93 0.94 0.95 0.96 0.97	3 4 5	1.40 1.42 1.43 1.44 1.45	6 7 8 9 39.0	1.92 1.93 1.94 1.95 1.96	1 2 3 4 5	2.50 2.51 2.52 2.54 2.55
3 4 5	0.05 0.06 0.08 0.09 0.10	6 7 8 9 26.0	0.52 0.53 0.54 0.55 0.57	1 2 3 4 5	0.98 0.99 1.00 1.01 1.02	6 7 8 9 35.0	1.46 1.47 1.48 1.49 1.50	1 2 3 4 5	1.98 1.99 2.00 2.02 2.03	6 7 8 9 44.0	2.56 2.58 2.60 2.61 2.63
6 7 8 9 22.0	0.11 0.12 0.13 0.14 0.15	3 4 5	0.58 0.59 0.60 0.61 0.62	6 7 8 9 31.0	1.03 1.04 1.05 1.06 1.07	1 2 3 4 5	1.51 1.52 1.54 1.55 1.56	6 7 8 9 40.0	2.04 2.05 2.07 2.08 2.09	1 2 3 4 5	2.64 2.65 2.67 2.68 2.70
1 2 3 4 5	0.16 0.17 0.18 0.19 0.20	6 7 8 9 27.0	0.63 0.64 0.65 0.66 0.67	1 2 3 4 5	1.08 1.09 1.10 1.11 1.12	6 7 8 9 36.0	1.57 1.58 1.59 1.60 1.61	3 4 5	2.10 2.12 2.13 2.14 2.15	6 7 8 9 45.0	2.71 2.72 2.74 2.75 2.77
6 7 8 9 23.0	0.21 0.22 0.23 0.24 0.25	1 2 3 4 5	0.68 0.69 0.70 0.71 0.72	6 7 8 9 32.0	1.13 1.14 1.15 1.16 1.17	3 4 5	1.62 1.64 1.65 1.66 1.67	6 7 8 9 41.0	2.16 2.18 2.20 2.21 2.23	3 4 5	2.78 2.79 2.80 2.82 2.84
1 2 3 4 5	0.26 0.27 0.28 0.29 0.30	6 7 8 9 28.0	0.73 0.74 0.75 0.76 0.77	3 4 5	1.18 1.19 1.20 1.22 1.23	6 7 8 9 37.0	1.68 1.69 1.70 1.71 1.72	3 4 5	2.24 2.25 2.26 2.27 2.28	6 7 8 9 46.0	2.85 2.87 2.88 2.89 2.90
6 7 8 9 24. 0	0.31 0.32 0.33 0.34 0.36	. 3 . 4 . 5	0.78 0.79 0.80 0.81 0.82	6 7 8 9 33.0	1.24 1.25 1.26 1.27 1.28	1 2 3 4 5	1.73 1.75 1.76 1.78 1.79	6 7 8 9 42.0	2.30 2.32 2.33 2.34 2.35	3 4 5	2.92 2.93 2.94 2.96 2.98
1 2 3 4 5	0.37 0.38 0.39 0.40 0.41	6 7 8 9 29.0	0.83 0.84 0.85 0.86 0.87	1 2 3 4 5	1.29 1.30 1.31 1.32 1.34	6 7 8 9 38.0	1.80 1.81 1.82 1.84 1.85	3 4 5	2.37 2.38 2.39 2.40 2.41	6 7 8 9 47.0	3.00 3.01 3.02 3.03 3.05

PERCENTAGES OF FAT CORRESPONDING TO SCALE READINGS ON THE WOLLNY REFRACTOMETER—(Continued).

											
Scale Read- ing.	Per Cent Fat.										
47.0	3.05	50.5	3-59	54.0	4.18	57-5	4.78	61.0	5-44	64.5	6.14
I	3.06	6	3.60	1	4.20	6	4.80	1	5.46	6	6.16
2	3.08	7	3.61	2	4.22	7	4.82	2	5.48	7	6.18
3	3.10	8	3.63	3	4-23	8	4.84	3	5.50	8	6.20
4	3.12	9	3.64	4	4-25	9	4.86	4	5-52	, 9	6.22
5	3.14	51.0	3.66	5	4.26	58.ó	4.88	5	5-54	65.0	6.24
6	3-15	1	3.67	6	4.28	1	4-90	6	5.56	1	6.27
7 8	3.16	2	3.68	7 8	4.29	2	4-92	7 8	5.58	2	6.29
	3.17	3	3.70	11	4.31	3	4-94		5.60	3	6.31
. 9	3.18	4	3-72	9	4-33	4	4-95	_ 9	5.61	4	6.34
48.0	3.20	5	3-74	55.0	4-35	5	4-97	62.0	5.63	5	6.36
ī	3.21	6	3.76	1	4-37	6	4.98	I	5.65	6	6.38
2	3-23	7	3.78	2	4.38	7	5.00	2	5.66	7	6.40
3	3-25	8	3.80	3	4-40	8	5.02	3	5.68	8	6.42
4	3-27	9	3.82	4	4-42	9	5.04	4	5-70	. 9	6.44
5	3.28	52.0	3.84	5	4-43	59.0	5.06	5	5-72	66.0	6.46
6	3-30	1	3.85	6	4-44	1	5.08	6	5-74		
7	3.32	2	3.87	7	4.40	2	5.10	7	5.76		
8	3-33	3	3.89	8	4.48	3	5.11	8	5-78		
9	3-34	4	3.90	9	4-49	4	5.13	, 9	5.80		
49.0	3.36	5	3.92	56.ó	4-51	5	5.15	63.0	5.82		
I	3.38	6	3-93	1	4-53	6	5.17	· I	5.84		
2	3.40	7	3.95	2	4-55	7	5.19	, 2	5.86		
3	3.42	8	3-97	3	4-57	8	5.20	3	5.00		
4	3.43	9	3.99	4	4-59	₄₋ 9	5.22	: 4	5.90		
5	3-44	53.0	4.01	5	4.60	60.0	5-24	5	5.92		
6	3-45	1	4.03	6	4.61	1	5.26	6	5-94		
7	3.46	2	4.04	7	4.63	2	5.28	7	5.96	1	
8	3.48	3	4.06	8	4.65	3	5.30	. 8	5.98		
9	3.50	4	4.07	9	4.67	4	5-32	, 9	6.00		
50.0	3.51	5	4.09	57.0	4.69	5	5-34	64.0	6.02		
1	3-53	6	4.10	r	4.71	6	5-36	' I	6.04		
2	3-55	7	4.12	2	4-73	7	5.38	2	6.07		l
3	3-56	8	4.14	3	4-75	8	5.40	, 3	6.09		
4	3-57	9	4.16	4	4.76	9	5-42	4	6.12		
5	3.59	54.0	4.18	5	4.78	61.0	5-44	5	6.14	1	
	ı ,	I	1 1	I.			'	1		11	I

The following table is of use for those who wish to employ the Wollny method, but have the Abbé refractometer instead of the milk-fat refractometer.

INDICES OF REFRACTION (n_D) CORRESPONDING TO SCALE READINGS OF THE WOLLNY MILK-FAT REFRACTOMETER.

Refrac- tive Index,				Fourt	h Decima	l of n _D .				
Index.	0	1	2	3	4	5	6	7	8	9
		,		Sea	de Readi	ngs.				,
1.333			0.0	0.1	0.2	0.3	0.4	0.5	0.5	0.6
1.334	0.7	0.8	0.9	1.0	1.1	1.2	1.3	1.4	1.5	1.6
1.335	1.7	1.8	1.9	2.0	2.I	2.1	2.2	2.3	2.4	2.5
1.336	2.8	2.7	2.8	2.9	3.0	3.1	3.2	3-3	3-4	3-5
1.337	3.6	3.7	3-7	3.8	3-9	4.0	4.I	4.2	4 - 3	4-4
1.338	4-5	4.6	4-7	4.8	4.9	5.0	5.I	5.2	5.3	5-4
1.339	5-5	5.6	5.7	5.8	5-9	6.0	6.1	6.2	6.3	6.4
1.340	6.5	6.6	6.7	6.8	6.9	6.9	7.0	7.1	7.2	7-3
1.341	7.4	7.5	7.6	7-7	7.8	7-9	8.0	8.1	8.2	8.3
1.342	8.4	8.5	8.6	8.7	8.8	8.9	9.0	9.1	9.2	9.3
1.343	9-4	9-5	9.6	9.7	9.8	9.9	10.0	10.1	10.2	10.3
1.344	10.4	10.5	10.6	10.7	10.8	10.9	11.0	11.1	11.2	11.3
1.345	11.4	11.5	11.5	11.6	11.7	11.8	11.9	12.0	12.1	12.2
1.346	12.3	12.4	12.5	12.6	12.7	12.8	12.9	13.0	13.1	13.2
1-347	13.3	13.4	13.5	13.6	13.7	13.8	13.9	14.0	14.1	14.2
1.348	14.3	14.4	14.5	14.6	14.7	14.8	14.9	15.0	15.1	15.2
1-349	15.3	15-4	15-5	15.6	15-7	15.8	15.9	16.0	16.1	16.2
1.350	16.3	16.4	16.5	16.6	16.7	16.8	16.9	17.0	17.1	17.2
1.351	17.3	17.4	17.5	17.6	17.7	17.8	17.9	18.0	ľ	18.2
1.352	18.3	18.4	18.5	18.6	18.7	18.8	18.9	19.0	19.1	19.2
. 1-353	19.3	19.4	19.5	19.6	19.7	19.8	19.9	20.0	20.1	20.2
1.354	20.3	20.4	20.5	20.6	20.7	20.8	20.9	21.0	21.1	21.2
1.355	21.3	21.4	21.5	21.6	21.7	21.8	21.9	22.0	22.1	22.2
1.356	22.3	22.4	22.5	22.6	22.7	22.8	22.9	23.0	23.1	23.2
1-357	23.3	23.4	23.5	23.6	23.7	23.8	23.9	24.0	24.I	24.2
1.358	-	24.4	24.5	24.6	24.7	24.8	24.9	25.0	25.1	25.2
1.359	25.3	25-4	25-5	25.6	25-7	25.8	25.9	26.0	26.1	20.2
1.360		26.4		26.6	26.7	26.8	26.9		27.1	27.3
1.361	27.4	27.5	27.6	27-7	27.8	27.9	28.0	28.1	28.2	28.3
1.362	28.4	28.5	28.6	28.7	28.8	28.9	29.0	29.1	29.2	29.3
1.363		29.5	29.6	29.7	29.8	29.9	30.0	30.1	30.2	30.3
1.364	30.4	30.5	30.6	30.7	30.8	31.0	31.1	31.2	31.3	31.4
1.365	31.5	31.6	31.7	31.8	31.9	32.0	32.1	32.2	32.3	32.4
1.366	32.5	32.7	32.8	32.9	33.0	33-1	33.2	33-3	33-4	33.5
1.367	33.6	33.7	33.8	33.9	34.0	34.2	34-3	34-4	34-5	34.6
1.368	34·7 35·7	34.8 35.8	34.9 36.0	35.0 36.1	35.1 36.2	35.2 36.3	35·3 36.4	35-4 36.5	35·5 36.6	35.6
										1
1.370	36.8	36.9	37.0	37.1	37.2	37-3	37-4	37.6	37.7	37.8
1.371	37.9	38.0	38.1	38.2	38.3	38.4	38.5	38.6	38.7	38.8
1.372	38.9 40.0	39.0 40.1	39.2 40.2	39-3	39-4	39.5	39.6	39.7	39.8	39.9
1.373	41.1	41.2		40.3	40.4	40.5	40.7		40.9	41.0
1.374	42.2	42.3	41.3	41.4	41.5	41.6	41.8 42.8	41.9 42.9	42.0	43.1
1.376	43.2	43.3	43.4	43.6		43.8	43.9	44.0	43.0 44.1	44.2
1.377	44-3	43.3	44.6	44-7	43-7 44.8	44.9	45.0	45.I	45.2	45.3
1.378	45.4	45.6	45-7	45.8	45-9	46.0	46.1	46.2	46.3	46.4
1.379	46.6	46.7	46.8	46.9	47.0	47.1	47.2	47.3	47-4	47.6
317	1 75.0	75.7		٠,٠,٠	7,	7′''	7,	77.3	7/**	17,.0

INDICES OF REFRACTION (n_D) CORRESPONDING TO SCALE READINGS OF THE WOLLNY MILK-FAT REFRACTOMETER—(Continued).

Refrac-	Fourth Decimal of n _D .											
Index, n _D .	0	1	2	3	4	5	6	7	8	9		
				,	Scale R	eadings.	,			-		
1.380	47-7	47.8	47-9	48.0	48.1	48.2	48.3	48.4	48.6	48.		
1.381	48.8	48.9	49.0	49.1	49.2	49-3	49-4	49.6	49-7	49.		
1.382	49-9	50.0	50.1	50.2	50.3	50.4	50.6	50.7	50.8	50.		
1.383	51.0	51.1	51.2	51.3	51.4	51.6	51.7	51.8	51.9	52.		
1.384	52.1	52.2	52.3	52-4	52.6	52.7	52.8	52.9	53.0	53-		
1.385	53-2	53.3	53.4	53.6	53-7	53.8	53-9	54.0	54.1	54.		
1.386	54-3	54-4	54.6	54-7	54.8	.54-9	55.0	55.1	55.2	55.		
1.387	55.4	55.6	55-7	55.8	55-9	56.0	56. r	56.2	50.3	50.		
1.388	56.6	56.7	56.8	56.9	57.1	57.2	57-3	57.4	57.6	57.		
1.389	57.8	57-9	58.0	58.1	58.2	58.3	58.4	58.6	58.7	58.		
1.390	58.9	59.0	59.1	.59.2	59.4	59-5	59.6	59.8	59-9	60.		
1.391	60.1	60.2	60.3	60.4	60.6	60.7	60.8	60.9	61.0	61.		
1.392	61.3	61.4	61.5	61.6	61.8	61.9	62.0	62.1	62.2	62.		
1.393	62.4	62.6	62.7	62.8	62.9	63.0	63.2	63.3	63.4	63.		
1.394	63.6	63.8 65.0	63.9	64.0	64.1	64.2	64.4	64.5	64.6	64.		
1.395	64.8 66.0	66.2	65.1 66.3	65.2	65.3	65.4	65.6	65.7	65.8	65.		
1.396	67.2	67.4	67.5	66.4	66.5	66.6 67.8	66.8	66.9 68.1	67.0 68.2	68.		
1.397	68.4	68.6	68.7	68.8	67.7 68.9	69.0	67.9			69.		
1.398	69.6	69.8	69.9	70.0	70.1	70.2	69.1 70.4	69.3 70.5	69.4 70.6	70.		
1,399	G.G	09.0	٠9.9	70.0	70.1	70.2	70.4	70.5	70.0	/0.		
1.400	70.9	71.0	71.1	71.2	71.4	71.5	71.6	71.8	71.9	72.		
1.401	72.1	72.2	72.4	72.5	72.6	72.8	72.9	73.0	73.1	73-		
1.402	73-4	73-5	73.6	73.8	73-9	74.0	74-1	74.2	74 - 4	74-		
1.403	74.6	74.8	74.9	75.0	7.5 - I	75-2	75.4	75.5	75.6	75-		
1.404	75-9	76.0	76.1	76.2	76.4	76.5	76.6	76.8	76.9	77.		
1.405	77.1	77.2	77-4	77.5	77-7	77.8	77-9	78.1	78.2	78.		
1.406	78.5	78.6	78.7	78.8	79.0	79.1	79.2	79-4	79-5	79.		
1.407	79.8	79.9	80.0	80.1	80.2	80.4	80.5	80.6	80.8	80.		
1.408	81.0	81.1	81.2	81.4	81.5	81.6	81.7	81.9	82.0	82.		
1.409	82.3	82.4	82.5	82.6	82.8	82.9	83.0	83.2	83.3	83.		
1.410	83.6	83.7	83.8	84.0	84.1	84.2	84.4	84.5	84.6	84.		
1.411	84.9	85.0	85.2	85.3	85.4	85.5	85.6	85.7	85.9	86.		
1.412	86.2	86.3	86.5	86.6	86.7	86.9	87.0	87.1	87.3	87.		
1.413	87.5	87.7	87.8	87.9	88.1	88.2	88.3	88.5	88.6	88.		
1.414	88.9	89.0	89.1	89.3	89.4	89.6	89.7	89.9	90.0	90.		
1.415	90.2	90.4	90.5	90.6	90.8	90.9	91.0	91.2	91.3	91.		
1.416	91.6	91.7	91.9	92.0	92.1	92.3	92.4	92.5	92.7	92.		
1.417	92.9	93.1	93.2	93-3	93.5	93.6	93.8	93-9	94.0	94-		
1.418 1.419	94·3 95·7	94·4 95.8	94.6 96.0	94.7 96.1	94.8 96.3	95.0 96.4	95.1 96.6	95·3 96.7	95·4 96.8	95 · 97 ·		
1.420	97.1	07.2	07.4	97.6	07.7	97.8	98.0	98.1	98.3	08.		
1.420	98.5	97·3 98·7	97·4 98.8	99.0	97·7 99.1	99.3	99.4	99.5	99.7	99.		
1.421	100.	90.7	90.0	99.0	99.1	99.3	yy•4	99.3	99.7	المح		

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DETERMINATION OF PROTEINS.—For determination of the total nitrogen in milk, 5 cc. are measured direct into a Kjeldahl digestion-flask, or a known weight from a weighing-bottle may be used, and the regular Gunning method is employed as described on page 69, proceeding with the digestion at once without evaporation.

The total nitrogen, multiplied by 6.38, gives the total proteins. By many the old factor of 6.25 is still employed, but in view of the fact that both casein and albumin have been found to contain 15.7% of nitrogen, there would seem to be the best reasons for employing 6.38 as a factor $\left(\frac{100}{15.7}\right)$.

Ritthausen's Method.—Ten grams of milk are measured into a beaker and diluted with water to about 100 cc. Five cc. of a solution of copper sulphate (strength of Fehling's copper solution, 34.64 grams CuSO, in 500 cc. of water) are added and the mixture stirred. A solution of sodium hydroxide (25 grams to the liter) is added cautiously a little at a time, till the liquid is nearly, but not quite neutral, avoiding an excess of alkali, as this would prevent the complete precipitation of the proteins. Allow the precipitate to settle, and pour off the supernatant liquid through a weighed filter, previously dried at 130° C. Wash a number of times by decantation. and transfer the precipitate to the filter, being careful to remove the portions adhering to the sides of the beaker with a rubber-tipped rod. thoroughly with water, and drain dry, after which the precipitate is washed with strong alcohol, dried, extracted with ether, preferably in a Soxhlet extractor, and then transferred on the filter to the oven, dried at 130° C., and weighed. The filter and precipitate are then burnt to an ash in a porcelain crucible, and the weight of the residue subtracted from the first weight gives that of the proteins.

Richmond* recommends modifying this process to the extent of neutralizing the milk, using phenolphthalein as an indicator, before adding the copper sulphate solution, and using only 2.5 cc. of the latter.

Determination of Casein.—Official Method of the A. O. A. C.—Ten grams of the milk are placed in a beaker, and made up with water to about 100 cc. at 40° to 42° C. One and one-half cc. of a 10% solution (by weight) of acetic acid are added, the mixture stirred, warmed to the above temperature, and allowed to stand for from three to five minutes, till a flocculent precipitate separates, leaving a clear supernatant liquid. Decant

^{*} Dairy Chem., p. 107.

upon a filter, wash with cold water two or three times by decantation, and finally transfer the whole of the precipitate to the filter, and, after filtering, wash two or three times. The filtrate should be clear or nearly so. If not, it can generally be made so by repeated filtrations, and the washing done afterwards. The filter containing the washed precipitate is transferred to the Kjeldahl digestion-flask and the nitrogen obtained by the Gunning process. $N \times 6.38 =$ casein.

Determination of Albumin.—Optional Methods of the A. O. A. C.—To the filtrate from the direct determination of casein by the acetic acid method as described in the preceding section, exactly neutralized with sodium hydroxide, 0.3 cc. of a 10% solution of acetic acid is added, and the mixture is boiled till the albumin is completely precipitated. The precipitate is collected on a filter and washed, the nitrogen being determined in the precipitate, and the factor 6.38 used in calculating the albumin therefrom.

Leffman and Beam's Modified Method for Albumin and Casein.—Owing to the tedious processes of washing and filtering incidental to the above methods for determining casein, the following is suggested. Twenty cc. of the milk are mixed with saturated magnesium sulphate solution, and the mixture saturated with the powdered salt. The whole is then washed into a graduate with a little of the saturated solution, and the precipitate allowed to settle, leaving a clear supernatant layer. The volume of the mixture in the graduate is read, and as much as possible of the clear portion is withdrawn by a pipette and filtered.

An aliquot part of the filtrate is then taken, and the albumin is precipitated from it by a solution of tannin, after which the precipitate is washed in a filter and the nitrogen determined therein. $N \times 6.38$ =albumin.

The casein is calculated by difference between the total proteins and the albumin.

Determination of Nitrogen as Caseoses, Amido-compounds, Peptones, and Ammonia.—Van Slyke * proceeds as follows: The filtrate from the determination of the albumin, as above, is heated to 70° C., 1 cc. of 50% sulphuric acid is first added, and afterwards chemically pure zinc sulphate to saturation. The mixture is allowed to stand at 70° until the caseoses separate out and settle. Cool, filter, wash with a saturated zinc sulphate solution slightly acidified with sulphuric acid, and determine the nitrogen of the caseoses in the precipitate.

^{*} N. Y. Exp. Station, Bul. 215, p. 102.

For Amido-compounds and Ammonia treat 50 grams of the milk in a 250-cc. graduated flask with 1 gram sodium chloride and a 12% solution of tannin, added drop by drop till no further precipitate is formed. Dilute to the 250-cc. mark, shake, and filter. Determine the nitrogen in 50 cc. of the filtrate, the result being the combined nitrogen of the amido-compounds and ammonia.

Distil with magnesium oxide 100 cc. of the filtrate from the tannin salt solution, receiving the distillate in a standardized acid, and titrating in the usual way for the ammonia.

Calculate the nitrogen of the *peptones* by subtracting from the total nitrogen that due to all other forms.

Van Slyke has furnished the following unpublished analysis of a sample of milk three months old, kept under antiseptic conditions by chloroform.

Per Cent	Per Cent	Per Cent N as Paranuclein, Caseoses, and Peptones.	Per Cent
Total N.	Sol. Nitrogen.		N as Amides.
0.561	0.099	0.074	0.025

DETERMINATION OF MILK SUGAR.—If a polariscope is available, the sugar of milk can most readily and conveniently be determined by optical methods. In the absence of a polariscope, the reducing power of milk sugar on copper salts may be utilized quite accurately in determining the sugar, using either volumetric or gravimetric methods as desired.

Determination by Optical Methods.—1. Reagents.—Acid Nitrate of Mercury.—This solution is prepared by dissolving metallic mercury in twice its weight of nitric acid of specific gravity 1.42, and adding to the solution an equal volume of water. One cc. of this reagent will be found sufficient to precipitate the proteins and fat completely from 65 grams of milk, but if more is employed the result of the analysis is not affected.

Mercuric Iodide Solution.—33.2 grams of potassium iodide are mixed with 13.5 grams of mercuric chloride, 20 cc. of acetic acid, and 640 cc. of water.

Subacetate of Lead Solution, U.S. P. See p. 586.

Notes.—For the Laurent polariscope, in which the normal weight for sucrose is 16.19 grams, the corresponding normal weight for lactose is 20.496, while for the Soleil-Ventzke instrument, in which the su-

crose normal weight is 26.048 grams, the corresponding lactose normal weight is 32.075.*

It is customary to employ three times the normal weight of milk in the case of the Laurent instrument (viz., 61.48 grams) and twice the normal weight in the case of the Soleil-Ventzke (viz., 65.95 grams).

As it is more convenient to measure the milk than to weigh it, and as the volume varies with the specific gravity, the following table is useful, showing the quantity to be measured in any case, having first determined the specific gravity.

	Volume of Milk to be Used.						
Specific Gravity.	For Polariscopes of which the Sucrose Normal Weight is 16.19 Grams.	For Polariscopes of which the Sucrose Normal Weight is 26.048 Grams.					
1.024	60.0 cc.	64.4 cc.					
1.026	59.9 cc.	64.3 cc.					
1.028	59.8 cc.	64. 15 cc.					
1.030	59.7 cc.	64.0 cc.					
1.032	59.6 cc.	63.9 cc.					
1.034	59.5 cc.	63.8 cc.					
1.035	59.35 cc.	63.7 cc.					

For ordinary work it is sufficiently close to have a pipette graduated to deliver 59.7 cc. if the Laurent instrument is used, and 64 cc. for the Soleil-Ventzke.

2. Process. — Measure as above, the equivalent of 61.48 grams of the milk for the Laurent, or 65.95 grams for the Soleil-Ventzke, instrument into a 100-cc. graduated flask, add, in order to clarify, 2 cc. of acid nitrate of mercury solution, or 30 cc. of mercuric iodide solution, or 10 cc. of lead subacetate solution. Shake gently and fill to the mark with water. Then add from a pipette enough water to make up for the volume of the precipitated proteins and fat, insuring 100 cc. of sugar solution. If the Laurent instrument is used, the amount added as prescribed by the A. O. A. C. is 2.4 cc., and with the Soleil-Ventzke 2.6 cc. The contents of the flask are then shaken and poured upon a dry filter. The filtrate, which should be perfectly clear, is polarized in a 200-mm. tube, and the reading, divided by 3 for the Laurent and by 2 for the Soleil-Ventzke, gives the percentage of lactose directly.

Allowance for the Volume of the Precipitate.—This of course varies

^{*} $[\alpha]_D$ for lactose= 52.53, $[\alpha]_D$ for sucrose=66.5, hence for the Laurent instrument 52.53: 66.5:: 16.19: 20.496, and for the Soleil-Ventzke instrument 52.53: 66.5:: 26.048: 32.975.

with the content in proteins, and fat, and while the above allowance gives in most cases sufficiently close results, it is not exact. Leffman and Beam * advise that the amount of water to be added above 100 cc. be calculated in each case from the percentage of proteins and fat previously found by analysis, multiplying the actual weight of the fat in grams in the sample taken by 1.075, and the weight of proteins by 0.8, the sum of the two results being the volume in cubic centimeters occupied by the precipitate.

All the calculations are avoided by employing the double-dilution method, which is to be recommended when very particular results are required.

Wiley and Ewell's Double-dilution Method.†—Two flasks are employed graduated at 100 and 200 cc. respectively, into each of which are introduced 65.95 grams of milk, if the Soleil-Ventzke instrument is used (or 61.48 grams in case the Laurent is used) and 4 cc. of the mercuric nitrate solution are added, both flasks being filled to the mark and shaken. The contents are filtered and the polarization is made in each case in a 400-mm. tube.

The second reading (that of the more dilute solution) is multiplied by 2, and the product subtracted from the first reading; the remainder is then multiplied by 2, and the product subtracted from the first reading (that of the stronger or 100 cc. solution). The result is the corrected reading, which, divided by 4, gives the exact per cent of milk sugar in the sample. This method depends on the fact that within ordinary limits the polarizations of two solutions of the same substance are inversely proportional to their volumes.

DETERMINATION OF MILK SUGAR BY FEHLING'S SOLUTION.—Twenty-five grams of the milk (24.2 cc.) are transferred to a 250-cc. flask, 0.5 cc. of a 30% solution of acetic acid are added and the contents well shaken. After standing for a few minutes, about 100 cc. of boiling water are run in, the contents again shaken, 25 cc. of alumina cream are next added, the flask shaken once more, and set aside for at least ten minutes. The supernatant liquid is then poured upon a previously wetted ribbed filter, and finally the whole contents of the flask are brought thereon, and the filtrate and washings made up to 250 cc. The filtrate must be perfectly clear. The milk sugar in a solution thus precipitated would ordinarily not exceed $\frac{1}{2}$ of 1 per cent.

^{*} Milk and Milk Products, p. 38.

[†] Wiley's Agricultural Analysis, p. 278; Analyst, 21, 1896, p. 182.

Volumetric Fehling Process.—From a burette containing the clear milk sugar solution above prepared, run a measured volume into the boiling Fehling liquor containing 5 cc. each of copper and alkali solution till sufficient has been introduced to completely reduce the copper, conducting the operation in the manner described in detail on page 591,

As 0.067 gram of milk sugar will reduce 10 cc. of Fehling solution (see p. 593), it follows that the number of cubic centimeters of sugar-containing solution required for making the test (using preferably the average of several determinations) will contain 0.067 gram of milk sugar, from which the percentage is readily computed. Thus if 16 cc. of the milk sugar solution are necessary to reduce the copper, then 16 cc. contain 0.067 gram milk sugar.

and 1.6 grams milk contain 0.067 gram milk sugar. Therefore the sample contains $\frac{.067 \times 100}{1.6} = 4.19\%$.

Gravimetric Fehling Processes.—O'Sullivan-Defren Method.—Twenty-five cc. of the above milk sugar solution are added to the hot mixture of 15 cc. each of Fehling copper and alkali solutions and 50 cc. water, prepared as directed on page 591 and the test carried out in accordance with the details there described. The weight of the cupric oxide, CuO, as formed, may be roughly calculated to anhydrous milk sugar by multiplying by 0.6024.

For more accurate results, however, the Defren table, page 595, should be used.

Soxhlet's Method.*—Twenty-five cc. of milk are diluted with 400 cc. of water in a half-liter graduated flask and 10 cc. of Fehling's copper solution are added. Then 8.8 cc. of half-normal sodium hydroxide are run in, or a sufficient quantity to nearly but not quite neutralize, the solution being still slightly acid. The flask is filled to the mark, shaken, and the contents filtered, using a dry filter.

One hundred cc. of the filtrate are added to 50 cc. of the mixed Fehling solution, which is boiled briskly in a beaker (using 25 cc. each of the copper and alkali solution). After boiling for six minutes, filter rapidly through a Gooch crucible provided with a layer of asbestos as described on page 594, and wash with boiling water till free from alkali. The asbestos

^{*} U. S. Dept. of Agric., Bur. of Chem., Bull. 46, p. 41; Bul. 107 (rev.), p. 119.

film with the adhering cuprous oxide is washed into a beaker by hot dilute nitric acid, and after complete solution of the copper is assured, it is again filtered and washed with hot water till a clean solution containing all the copper is obtained. Add 10 cc. of dilute sulphuric acid (containing 200 cc. of sulphuric acid, specific gravity 1.84 per liter) and evaporate on the steambath till the copper has largely crystallized, then carefully continue the heating over a hot plate till the nitric acid is driven out, as evidenced by the white fumes of sulphuric. Add 8 or 10 drops nitric acid (specific gravity 1.42) and rinse into a very clean tared platinum dish of about 100 cc. capacity, in which the copper is deposited by electrolysis. See page 608.

The weight of milk sugar is determined from that of copper found, from the table on page 152.

If the apparatus for the determination of the copper by the electrolytic method is not at hand, the cuprous oxide may be weighed directly in the Gooch crucible. In order to facilitate drying, it should be washed successively with 10 cc. of alcohol, and 10 cc. of ether, after which it is dried thirty minutes in a water-oven at 100° C., cooled, and weighed. The weight of copper is obtained from the weight of the cuprous oxide by the use of the factor 0.8883.

Munson and Walker Method.—The milk sugar solution is prepared as in Soxhlet's method. For details as to the copper reduction process see page 598.

Relation between Specific Gravity, Fat, and Total Solids of Milk.— The close relationship existing between these factors has long been known, and many formulæ have been devised, whereby, if two of them are known, the third may be computed with considerable approach to accuracy. The specific gravity and the fat are very readily determined by any dairyman, by the aid of a lactometer and the Babcock apparatus. The total solids are ascertained with more difficulty, since the use of more involved and costly apparatus is necessary, besides considerable technical skill. It is therefore common for producers to calculate the total solids from the fat and specific gravity, using one of the many tables prepared for the purpose, based on some one of the best accepted formulæ. The total solids can thus be calculated to within two or three tenths of a per cent.

The two most commonly used formulæ for this purpose are those of Hehner and Richmond in England, and Babcock in the United States. Hehner and Richmond's formula is

$$T = 0.25S + 1.2F + 0.14$$

SOXHLET'S TABLE FOR THE DETERMINATION OF LACTOSE.*

Milli- grams of Cop- per.	Milli- grams of Lac- tose.								
100	71.6	161	117.1	221	162.7	281	209.1	341	256.5
101	72.4	162	117.9	222	163.4	282	209.9	342	257-4
102	73-1	163	118.6	223	164.2	283	210.7	343	258.2
103	73.8	164	119.4	224	164.9	284	211.5	344	259.0
104	74.6	165 166	120.2	225	165.7 166.4	285 286	212.3	345	259.8 260.6
105 106	75-3 76.1	167	120.9	227	167.2	287	213.1	346	261.4
107	76.8	168	121.7	228	167.9	288	214.7	347 348	262.3
108	77.6	160	123.2	220	168.6	280	215.5	349	263.1
100	78.3	170	123.9	230	169.4	290	216.3	350	263.9
110	79.0	171	124.7	231	170.1	201	217.1	351	264.7
III	79.8	172	125.5	232	170.9	292	217.9	352	265.5
112	8ó.5	173	126.2	233	171.6	293	218.7	353	266.3
113	81.3	174	127.0	234	172.4	294	219.5	354	267.2
114	82.0	175	127.8	235	173.1	295	220.3	355	268.0
115	82.7	176	128.5	236	173.9	296	221.1	356	268.8
116	83.5	177	129.3	237	174.6	297	221.9	357	269.6
117	84.2	178	130.1	238	175.4	298	222.7	358	270.4
118	85.0	179	130.8	239	176.2	299	223.5	359	271.2
119	85.7	181	131.6	240	176.9	300	224.4	360 361	272.1
120 121	86.4 87.2	182	132.4	241	177.7	301	225.2 225.9	362	272.9
121	87.9	183	133.1	243	179.3	303	225.9	363	273.7
123	88.7	184	134.7	244	180.1	304	227.5	364	274·5 275·3
124	89.4	185	135.4	245	180.8	305	228.3	365	276.2
125	90.1	186	136.2	246	181.6	306	220.I	366	277.I
126	90.0	187	137.0	247	182.4	307	229.8	367	277.9
127	ģ1.6	188	137.7	248	183.2	308	230.6	368	278.8
128	92.4	189	138.5	249	184.0	309	231.4	369	279.6
129	93.1	190	139.3	250	184.8	310	232.2	370	280.5
130	93.8	191	140.0	251	185.5	311	232.9	371	281.4
131	94.6	192	140.8	252	186.3	312	233-7	1 3/-	282.2
132	95-3	193	141.6	253	187.1	313	234.5	373	283.1
133	96.1	194	142.3	254	187.9	314	235-3	374	283.9
134	96.9	195	143.1	255	188.7	315	236.1 236.8	375	284.8
135	97.6	196	143.9 144.6	256 257	189.4	316 317	237.6	376	285.7 286.5
136 137	90.3	198	145.4	258	191.0	318	238.4	377 378	287.4
138	99.8	199	146.2	259	191.8	319	239.2	379	288.2
139	100.5	200	146.9	260	192.5	320	240.0	380	280.1
140	101.3	201	147-7	261	193.3	321	240.7	381	280.0
141	102.0	202	148.5	262	194.1	322	241.5	382	29ó. Ś
142	102.8	203	149.2	263	194.9	323	242.3	383	291.7
143	103.5	204	150.0	264	195.7	324	243.I	384	292.5
144	104.3	205	150.7	265	196.4	325	243.9	385	293.4
145	105.1	206	151.5	266	197.2	326	244.6	386	294.2
146	105.8	207	152.2	267	198.0	327	245.4	387	295.1
147	106.6	208	153.0	268	198.8	328	246.2	388	296.0
148	107.3	209	153-7	269	199.5	329	247.0	389	296.8
149	108.1	210	154.5	270	200.3	330	247.7	390	297.7
150	100.6	211	155.2	271	201.1	331 332	249.2	391	298.5
151 152	110.3	213	156.7	273	202.7	333	250.0	392 393	300.3
153	111.1	214	157-5	274	203.5	334	250.8	393	301.1
154	111.9	215	158.2	275	204.3	335	251.6	395	302.0
155	112.6	216	159.0	276	205.1	336	252.5	396	302.8
156	113.4	217	159.7	277	205.9	337	253.3	397	303.7
157	114.1	218	160.4	278	206.7	338	254.1	398	304.6
158	114.9	219	161.2	279	207.5	339	254.9	399	305.4
159	115.6	220	161.9	280	208.3	340	255.7	400	306.3
160	116.4	li .	!	11	L	11	1	II.	1

^{*} Wiley, Principles and Practice of Agricultural Analysis, Vol. III. pp. 163-165.

FIG. 49-KIChmond's Milk-scale.

where T is the per cent of total solids, S the lactometer reading, and F the fat. An ingenious instrument known as Richmond's milk-scale (Fig. 49) is useful in making the calculation, instead of employing either the formula or a table. This is constructed on the principle of the slide rule, and by its use the specific gravity may be corrected to the proper temperature, and the solids calculated from the fat and specific gravity.

Babcock's formula for solids not fat is as follows:

Solids not fat
$$= \left(\frac{100S - FS}{100 - 1.0753FS} - 1\right)(100 - F)2.5$$
,

S being the specific gravity, and F the percentage of fat. On this formula he has prepared a table * by means of which one may calculate solids not fat agreeing quite closely with results obtained by gravimetric analysis.† The table on page 154 has been recomputed and enlarged from that of Babcock, so as to express results in total solids rather than solids not fat.

Calculation of Proteins.—Van Slyke's \ddagger formula for calculating proteins (P) from the fat (F) is:

$$P = (F - 3) \times 0.4 + 2.8$$
.

Olsen § has devised the following formula for calculating proteins from total solids (TS):

$$P = TS - \frac{TS}{1.34}$$

Approximately 0.8 × proteins = casein.

The proteins being thus calculated, the sugar may be computed by difference. These calculations, while only approximate, give quite satisfactory results for normal, healthy milk, especially from herds.

Determination of Acidity.—While milk is still fresh, i.e., before it has begun to undergo factic fermentation, it will show an acid reaction, which is sometimes expressed in terms of factic acid. In view of the fact that

^{*} U. S. Dept. of Agric., Div. of Chem., Bul. 47, p. 123; Bul. 107 (rev.) p. 225.

[†] For approximate work Babcock has suggested the following simplified formulæ: Solids not fat=0.25G+0.2F and total solids=0.25G+1.2F, G being the lactometer reading and F the fat.

^{\$} Jour. Am. Chem. Soc. 30, 1908, p 1182.

[§] Jour. Ind. and Eng. Chem., 1, 1909, p. 253.

TABLE SHOWING PER CENT OF TOTAL SOLIDS IN MILK CORRESPONDING TO QUEVENNE LACTOMETER READINGS* AND PER CENT OF FAT.†

Cent -		
0.0		Lactometer Reading at 15.5° C.
0.2 0.3 10 7 7.13 7.37 7.63 7.65 7.66 7.67 8.12 8.37 8.61 8.47 0.01 7.66 8.11 8.10 8.01 8.56 0.01 9.36 0.00 9.36 0.0	of Pat.	28
0.3 0.4 11.7 36 7 74 7 74 7 76 8 8.44 8.40 8 74 8.00 9 24 8.00 9 24 9 7 74 7 74 7 70 8 8.44 8.40 8 74 8.00 9 24 9 7 8 7 74 7 74 7 70 8 8.41 8.40 8 74 8.00 9 24 9 7 7 8 7 8 7 7 8 7 7 8 7 8 8 8 8 8 8 8		75 7 00 7 25 7 50 7 75 8 00 8 25 8 50 8.75 9.00
0. t 33	0.2	10 7 24 7 49 7 74 7 90 8 24 8 40 8 74 8 90 9 2
0.6 0.6 0.7 17 17 18 17 18 18 18 18 18 18 18 18 18 18 18 18 18	_	
10	0.5	1 351 7 60 7.85 8.10 8 35 8 60 8.85 9.10 p 35 9.66
0.8 71 700 8		17 7 72 7 97 8 22 8.47 8.72 8 97 9.22 9.47 9 7: 50 7.84 8 00 8.34 8 50 8.84 9.00 9 34 9 50 9 8.
1.2 1.2 1.3 1.4 1.3 1.4 1.3 1.4 1.5 1.5 1.5 1.6 1.6 1.7 1.6 1.7 1.7 1.7 1.7 1.7 1.7 1.7 1.8 1.8 1.8 1.8 1.8 1.8 1.8 1.8 1.8 1.8	0.8	77 7 90 8 31 8 40 8.71 8.96 9 31 9 40 9.72 9 90
1.3 1.4 1.3 1.4 1.8 1.6 1.6 1.7 1.6 1.7 1.8 1.8 1.8 1.8 1.9 1.9 1.7 1.8 1.9 1.9 1.9 1.9 1.9 1.9 1.9 1.9 1.9 1.9		25 8 20 8 45 8 70 8 95 9.20 9 45 9 70 9 95 10 20
1.4 1.4 1.8 1.8 1.8 1.8 1.8 1.8		
1.6 1.6 1.6 1.7 1.6 1.7 1.6 1.7 1.6 1.7 1.6 1.7 1.7 1.6 1.7 1.7 1.8 1.8 1.8 1.9 1.8 1.9 1.8 1.9 1.8 1.9 1.9 1.8 1.9 1.9 1.8 1.9 1.9 1.9 1.9 1.9 1.9 1.9 1.9 1.9 1.9		3 56 8 81 9.06 9 31 9 56 9.81 to 06 10.31 to 96
1.8 1.8 1.9 1.8 1.9 1.9 1.8 1.9 1.9 1.8 1.9 1.9 1.8 1.9 1.9 1.8 1.9 1.9 1.8 1.9 1.9 1.8 1.9 1.9 1.9 1.9 1.9 1.9 1.9 1.9 1.9 1.9	1.5	55 8 80 9 05 9 30 9 55 9 80 to 05 to 30 10 55 to 80
3.0		
3.0 2.1 3.7 3.0 3.3 3.4 2.5 3.5 3.6 3.6 3.7 3.7 3.7 3.7 3.7 3.7 3.7 3.7 3.7 3.7	1.8) 9.16 9.41 9.66 9.91 to.16 to.41 to 66 to 91,11,11
3.2 3.9 3.9 3.9 3.9 3.9 3.9 3.9 3.9 3.9 3.9		
1		17 0.52 9.77 10 03 10.27 10 52 10.78 11 03 11.28 11.3
3		[39 9 64 9 89 10 14 to 39 to 64 to 90 tr. 15 11 . 40 tr 6
2 7 8 60 8 87 0.12 0.37 0 62 0 87 10.12 10 37 10.02 10 87 11 12 11.38 17 6 311.88 12.18 2 7 8 87 4 8 99 0.24 0 99 0 74 0 90 0.74 10 90 17 41 10.90 11.74 11.50 11.75 12 00 12.28 2 8 8 8 0 11 0 36 0.61 0 86 10 11 10 36 10.11 11 13 71 10 21 11 87 12.12 13 37 10 21 11 87 12.12 13 37 10 21 11 87 12.12 13 37 10 21 11 87 12.12 13 37 10 21 11 87 12.12 13 37 10 21 11 87 12.12 13 37 10 21 11 87 12.12 13 37 10 21 11 87 12.12 13 37 10 21 11 87 12.12 13 37 10 21 11 87 12.12 13 37 10 21 11 87 12.12 13 37 10 21 11 87 12.12 13 37 10 21 11 87 12.12 13 13 13 13 13 13 13 13 13 13 13 13 13	3.4	53 9 88 to. 13 10 38 to 53 to 58 11.14 tr. 30 11.64 11.86
2 7 8 74 8 99 9.24 9 49 9 74 9 90 10.24 10 40 10.71 10 36 10.86 11.1121 37 11 0.11 18 71 12.12 12 37 2.98 8 86 9 11 9 36 9.61 9 86 10 11 10 36 10.51 10.86 11.1121 37 11 0.11 18 71 12.12 12 37 2.98 8 98 9 23 9.48 9 73 9 98 10.23 10 48 10.73 10.08 11.23 11.40 11.74 11 99 12 24 12.49 3.0		
3.0 0 10 9 35 0 60 0 85 10 10 10 35 10 60 10 85 11.10 11.36 11 61 11.86 12 11 12 36 12.61 3.1 0 9.22 9.47 0.72 0.07 10.22 10 47 10.72 10.07 11.23 11.48 11.73 11.05 12.23 12.48 12.73 12.63 3.2 0 34 0 50 0 85 10 05 10 31 0 50 10.55 11.00 11.35 11.60 11.85 12.23 12.48 12.73 12.63 3.3 0 46 0 77 10.05 10.22 10 40 10.71 10 00 11.21 (1.47 11.72 11.05 11.85 12.23 12.48 12.73 12.63 3.4 0 68 9 8 110.65 10 310 58 10.85 11.00 11.34 11.50 11.83 12.00 12.24 48 12.73 12.63 3.5 0 70 0.05 10.20 10 45 10 70 10 05 11.21 11.40 11.71 11 00 12 21 12 40 12 72 12 07 13 28 3.5 0 98 10 0.5 10 31 10 58 10.55 10.55 11.55 11.55 11.50 11.35 12.50 12.35 12.35		8 74 8 90 0.24 0 40 0 74 9 90 10.24 20.40 10 74 10 90 17 24 17.50 17.75 12 00 12.20
3.1		8 98 9 23 9.48 9 73 9 98 10.23 to 48 to 73 to 98 tt.23 11.49 tt.74 tt 99 12 24 12.40
3.3 9 46 9 71 0.06 to 21 to 40 to 71 to 00 tf.22 tf.47 tf.71 tf.70 tf.8 ff.7.10 tf.2 35 ff.2 61 ff.2 86 3.3 9 46 9 71 0.06 to 21 to 40 to 71 to 00 tf.22 tf.47 tf.7 tf.72 tf.67 tf.8 ff.7.10 tf.2 21 tf.48 ff.7 ff.2 22 tf.48 ff.7 ff.2 35 ff.2 68 3.4 9 58 9 51 to 08 to 33 to 58 to 08 tf.22 tf.46 tf.7 tf.7 tf.72 tf.67 tf.8 ff.7 tf.2 22 tf.48 ff.2 73 tf.2 98 3.5 9 70 0.05 to .20 to 45 to 70 to 09 tf.2 tf.47 tf.7 tf.7 tf.0 tf.2 21 tf.46 tf.7 tf.7 tf.2 tf.2 tf.2 tf.2 tf.2 tf.2 tf.2 tf.2		
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6.0 12.71 12.96 13.21 13.46 13.71 13.96 14.22 14.47 14.72 14 98 15 23 15.48 15.73 15.98 16.24		

The lactometer reading is expressed in whole numbers for convenience. The true specific gravity corresponding to a given lactometer reading is obtained by writing 1.0 before the lactometer reading. Thus, 1 026 is the specific gravity corresponding to lactometer reading 25, etc.

[†]An. Rep. Mass. State Board of Health, 1901, p. 445. (Analyst's Reprint, p. 25.)

the acidity of "sweet" milk is due partly to the presence of acid phosphates and partly to dissolved carbonic acid in the milk, and not to lactic acid, which is probably absent, a better plan is to express the acidity in terms of the number of cubic centimeters of tenth-normal alkali necessary to neutralize a given quantity of the milk, either 25 or 50 cc., using phenolphthalein as an indicator.

If it is desired to calculate the acidity in terms of lactic acid, multiply the number of cubic centimeters of tenth-normal alkali used by 0.897, and divide by the number of cubic centimeters of milk titrated, the result being the percentage of lactic acid.

Detection of Boiled Milk.—Storch's Method.*—Shake 5 cc. of the milk in a test-tube with one drop of a 2% solution of hydrogen peroxide and two drops of a 2% solution of paraphenylenediamin. If the milk has not been heated beyond 80° C., a dark violet color appears at once, but if it has been pasteurized or boiled, no color appears. Siegfeld and Samson † find that addition of two drops of formalin (1:1) to each 100 cc. of milk previous to boiling causes it to react similar to raw milk.

MODIFIED MILK.

A comparison of the composition of cow's milk and human milk, as in the following table by Dr. Emmett Holt,‡ shows very marked differences.

	Woman's Milk, Average.	Cow's Milk, Average.
Fat	4.00	3.50
Sugar	7.∞	4.30 .
Proteins	1.50	4.00
Ash	0.20	0.70
Water	87.30	87.50

The per cent of fat in the two kinds of milk is nearly the same. There is, however, too little sugar and an excess of proteins and ash in the milk of the cow, assuming human milk as the ideal infant food, so that in basing a diet for infants on the basis of human milk considerable modification is necessary. Moreover, aside from the actual variation in the amount of ingredients, there are certain inherent differences in the character of the same ingredient, as found in the milk of the cow and in

^{*} Copenhagen Expt. Sta., 40th Rep. † Molk. Ztg., 21, 1907, p. 103. † "Infancy and Childhood."

human milk. The proteins of cow's milk are, for instance, found to be much more difficult of digestion than those of woman's milk, and the same is probably true of the fat. Aside from the mere statement of a few of these differences, it is obviously beyond the scope of this work to discuss this phase of the subject in detail, reference being made, however, to such books as Dr. T. M. Rotch's "Pediatrics," and "Infancy and Childhood" by Dr. Emmett Holt, for full particulars. So great has been the demand by physicians for "modified milk" for infant feeding, that laboratories for this exclusive purpose have been established in many of the larger cities, in which not only is milk prepared in accordance with certain fixed formulæ supposed to be adapted to average infants of varying age, but milk of any desired composition is prepared, in accordance with special prescriptions of physicians to apply to individual cases.

Methods and Ingredients.—The proteins and the ash in cow's milk are much higher than in human milk, and both are brought to the proper degree of reduction by diluting the milk with water. Milk sugar is increased by the addition of lactose, and the fat is increased or diminished by addition of cream or by skimming.

The dilution of cow's milk with a measured amount of water shows the following results on the proteins and ash:

	Cow's Milk.	Diluted Once.	Diluted Twice.	Diluted Three Times.	Diluted Four Times.
Proteins		Per cent. 2.00 0.35	Per cent. 1-33 0.23	Per cent. 1.00 0.18	Per cent. 0.80 0.14

The ingredients commonly employed for modifying milk are (1) cream, containing 16% of fat, (2) centrifugally skimmed milk, otherwise known as "separator milk" from which the fat has been removed, (3) milk sugar, or a standard solution of milk sugar of, say, 20% strength, and (4) lime water. Unusual care should be taken in the selection of the milk supply to insure cleanness, purity, and freshness, as well as in the care of utensils, etc., used in the laboratory, which should in all cases be scrupulously clean. Samples prepared in accordance with a given formula or formulæ are pasteurized in separate bottles, or, if desired, sterilized, and after stoppering with cotton are kept on ice.

Formulæ.—It is obviously impossible to establish formulæ universally applicable even to healthy infants, but the following may be regarded as typical formulæ, representing the composition of modified milk to suit the needs of an average growing infant during its first year:

Period.	Fat.	Proteins.	Sugar.
	Per cent.	Per cent.	Per cent.
Third to fourteenth day	2	0.6	6
Second to sixth week	2.5	0.8	6
Sixth to eleventh week	3	1.0	6
Eleventh week to fifth month	3-5	1.5	7
Fifth to ninth month	4	2	7
Ninth to twelfth month	3-5	2-5	3-5

Milk according to the above formulæ can be very simply prepared by the aid of a specially made graduate known as the "Materna" and shown in Fig. 50.

PREPARED MILK FOODS.

Milk Powder.—There are numerous brands of desiccated milk or milk powder on the market, sold in bulk and by the can, and largely used by bakers and manufacturers of milk chocolate. Many of these, purporting to contain all the ingredients of milk excepting water, have been found by the author to be pulverized dried skimmed milk. The following are analyses of whole milk, half-skim milk, and skim milk powders:



Fig. 50.—The "Materna" Graduate for Modifying Milk.

	Whole Milk, Powder.*	Half-skim Milk, Powder.*	Skim Milk, Powder.†
Moisture	. 3.62	5.01	8.16
Fat	. 26.75	15.26	1.73
Proteins $(N \times 6.25)$. 32.06	38.39	33.84
Milk sugar	. 31.90	34.67	49-35
Ash	- 5.67	6.67	6.87
	100.00	100.00	99.95

The fat in the skim milk powder corresponds to about 0.16% fat in the original milk.

Jensen ‡ states that the casein of dried milk no longer has the power

^{*} C. Huyge, Rev. gén. du Lait, 3, 1904, p. 400.

[†] Analysis by the author.

[‡] Molkerei Ztg., Berlin, 15, 1905, p. 565.

of swelling when mixed with water. To obviate this difficulty, Hatmaker adds to the milk 1 to 3% of sodium bicarbonate, and Elkenberg 2% of cane-sugar. A Swiss milk powder examined by Jensen contained an excess of sodium and a low acidity, indicating the addition of an alkaline sodium salt.

Artificial Albuminous Foods.—The albumin and casein of milk have furnished the basis of a variety of food preparations, some of which are intended for the use of invalids and people of weak digestion, and others, from their compactness, for travellers and campers. Among these foods are the following:

Nutrose.—This is a caseinate of sodium formed by the action of the alkali upon dried casein. It is soluble in water.

Eucasin is a caseinate of ammonium, a soluble powder somewhat similar to nutrose.

Plasmon.—This is a yellowish powder, prepared by treatment with sodium bicarbonate of the curd precipitated from skimmed milk. The compound is kneaded in an atmosphere of carbon dioxide, and reduced to a soluble powder.

The following analysis of plasmon was made by Woods and Merrill:*

Water.	Proteids.	Pat.	Carbohydrates.	Ash.	Fuel Value.	
8.5	75.0	0.2	8.9	7-4	2044	

Sanose.—This is also a powder, containing 80% of pure casein and 20% of albumose, obtained from the white of egg. The powder possesses a slight taste and an odor suggestive of milk. By briskly stirring the powder with water, an emulsion may be made much resembling milk, but on standing it soon breaks up.

Sanatogen is a grayish-white, tasteless powder, containing 95% of casein and 5% sodium glycero-phosphate. When treated with cold water it swells, forming on heating a milk-like emulsion.

Koumis is a stimulating beverage, prepared by allowing milk to undergo alcoholic, lactic, and proteolytic fermentations. The original koumis was made by the Tartar tribes of Asia from mare's milk, which contains more lactose than cow's milk, and apparently lends itself more readily to fermentation. Only a limited amount of koumis is now made from mare's milk, the milk chiefly used for this preparation being that of the cow, treated with yeast and sometimes added sugar. Koumis is a beverage much more commonly used in Europe than in America.

^{*} Maine Exp. Station, Bulletin 178, p. 101.

The following analyses were made by Vieth:*

	Water.	Alco- hol.	Fat.	Casein.	Albu-	Albu- min- oses.	Lactic Acid.	Sugar.	Ash.
Mare's milk	90.57	2.98 1.04 0.57	1.30 1.38 0.33	1.88	0.24 0.20 0.07	0.77		2.18	0.35 0.58 0.84

Kephir.—This is a fermented milk product similar to koumis, excepting that the fermentation is induced by a fungus known as kephir grains. The proteolytic fermentation is less pronounced in kephir than in koumis. König gives the following table as the mean of twenty-eight analyses:

Water.	Netro- gen.	Casein.	Albu- min.	Acid Albu- min.	Hemi- albumin.	Pep- tone.	Fat.	Lac- tose.	Lactic Acid.	Alco- hôl.	Ash.
91.21	3-49	2.53	0.36	0.21	0.21	0.039	1.44	2.41	1.02	0.75	0.68

ADULTERATION OF MILK.

Systems of Milk Inspection.—A typical method of general food inspection has already been outlined (see pp. 6 and 8), which may easily be modified to include the inspection of milk in connection with other foods, or to provide for a system of milk inspection exclusively. In the examination of such a perishable food as milk, it has not been found practicable for the analyst to reserve for the benefit of the defendant a sealed sample, as in the case of other foods, but experience has shown it had best be made the duty of the collector or inspector to give a sealed sample of milk to the dealer, when the latter requests it at the time of taking the sample. For this purpose the collector is provided with small bottles and sealing pharaphernalia, in addition to the tagged sample bottles or cans in which he collects the milk. The collector should use the same precautions for obtaining a perfectly fair representative sample as does the chemist in making the analysis, i.e., he should carefully pour the milk from the original container into an empty can or vessel and back again, before taking his sample.

Each sample is properly numbered by the collector in presence of the dealer, and the data as to the taking of the sample entered at once under the proper number in the collector's book. If a sealed sample is given,

^{*} Richmond Dairy Chemistry, p. 241 et seq.

it should bear the same number as the sample reserved for analysis, and a receipt should invariably be required from the dealer, as evidence that his request for a sealed sample has been complied with.

Milk Standards Fixed by Law.—In localities where a systematic form of milk inspection prevails, there is usually in force a statute fixing the legal standard for the total solids, and in many cases for the fat or for the solids exclusive of fat. In some states the statute is so drawn that any deviation from the legal standard constitutes an adulteration in the eye of the law, and hence the offender, who has such milk in his possession with intent to sell, is liable to the same fine as if he actually added water or a foreign substance to the milk.

In other states a distinction is made by the statute between milk that is simply below the legal standard of total solids, and milk containing actually added ingredients (water or otherwise), a much lighter fine being imposed for the former than for the latter offense. Where such a distinction prevails, it often becomes incumbent upon the analyst to show to the satisfaction of the court, in case of milk low in solids, whether or not the milk has been fraudulently watered after being drawn from the cow, it being well understood that cows may give milk below the standard.

Pure milk that is low in solids may owe its deficiency either to poor feeding, or to an inherent tendency on the part of the cow to give milk always of poor quality. Thus the Holstein cow, more than any other breed, is open to the charge of sometimes giving milk below the standard.* That the Holstein cow is a favorite with the producer is by no means strange, from the fact that no other breed can with moderate feeding be made to give so large a quantity of milk.

Wherever there is a statute fixing the standard for milk, it commonly provides also that the addition of any foreign substance whatsoever constitutes an adulteration.

U. S. Standards.†—Standard milk is the fresh, clean, lacteal secretion obtained by the complete milking of one or more perfectly healthy

^{*} This statement should not be taken as condemning the Holstein, for it is true that cows of this breed often give milk far above the standard. A large number of samples of milk of known purity from Holsteins analyzed by the writer have been found to be of excellent quality. It is a curious fact that among the samples of known purity analyzed by the Massachusetts Board of Health, both the lowest and highest total solids on record came from a Holstein cow; the lowest recorded total solids in a "known purity" milk being 9.96 per cent. (seventh annual report of Massachusetts State Board of Health, Lunacy, and Charity, p. 160), and the highest being 17.06 per cent. (twenty-second annual report of the Massachusetts State Board of Health, p. 405).

[†]U. S. Dept. of Agric., Off. of Sec., Circ. 19.

cows, properly fed and kept, excluding that obtained within fifteen days before and ten days after calving, and contains not less than 8.5% of solids not fat, nor less than 3.25% of milk-fat.

Standard Skim-milk is skim-milk containing not less than 9.25% of milk solids.

FORMS OF ADULTERATION.—Milk is ordinarily adulterated (1) by watering, (2) by skimming, (3) by both watering and skimming, and (4) by the addition of one or more foreign ingredients.

Watering and Skimming.—The fact that milk is found below the standard of total solids, while more often due to an excess of water, may also be due to a deficiency in fat. In one case the milk is commonly termed watered, and in the other skimmed, using the terms broadly and not necessarily meaning actual and fraudulent tampering with the milk. In a third case, and almost invariably fraudulently, both watering and skimming may be found to have been practiced on the same sample. The analyst judges which of these causes have produced a milk low in solids, by a careful study of the relation between the percentages of total solids, fat, and solids not fat.

If both the total solids and solids not fat are abnormally low, and the proportion of fat to solids not fat about the same as, or higher than, in a normal milk, it is generally safe to assume that the sample has been watered; if both the total solids and the fat are well below the standard, and the solids not fat nearly normal, then the milk has undoubtedly been skimmed; if, in the third place, the total solids and the solids not fat are proportionally reduced below the standard, while the ratio of fat to solids not fat is abnormally small, it is safe to adjudge the milk to be low by reason of both skimming and watering.

Milk of Known Purity.—It is difficult to place the minimum figure for total solids, below which a milk sample may safely be pronounced by the analyst as fraudulently watered after having been drawn from the cow. Nearly nine hundred samples of milk of known purity from various breeds of cow, milked in the presence of an inspector, have been analyzed in the Department of Food and Drug Inspection of the Massachusetts State Board of Health, extending over a period of fifteen years, and among these are many samples from Holstein cows. It is extremely rare that any of these known purity samples have been found with total solids as low as 11%, though there are instances where total solids have run as low as 10%.

It is safe to assume that in the few cases on record showing less than 10.75% of total solids, either there was something decidedly abnormal about the health of the cow, or, through some accident, the cow was only partially milked, it being a well-known fact that the last fraction of the milking includes the larger percentage of fat. (See page 128.)

It is therefore nearly always safe to condemn a milk standing below 10.75 as fraudulently watered, if at the same time it has a proportionately high per cent of fat.

The average total solids of 800 samples of milk of known purity analyzed by the Massachusetts Board up to and including the year 1890 amounted to about 131%.

It is rare indeed to find a herd of ten or more well-fed cows of mixed breeds in which the average milk of the herd falls below 12½% of solids.

The milk of forty-seven Holstein cows, examined in 1885, was found to contain an average of 12.51% of total solids, while the milk of eleven Jerseys examined in the same year averaged 14.02% of solids. These examples represent the two extremes commonly met with.

Variation in Standard.—In Massachusetts the law fixes a different standard for total solids in milk during the summer, or pasture-fed season, from that in force during the winter, or stall-fed period. From April to September inclusive the legal standard is 12% of total solids, of which 9% are solids not fat, and from October to March inclusive it is 13%, of which 9.3% are solids not fat. Bearing on the question of difference in normal quality of milk during the two periods, averages were taken of the milks collected by the corps of inspectors of the Massachusetts Board of Health during a month in each period, December and June being selected as most typical, and during these months all the samples were analyzed both for total solids and fat. The samples were taken from stores, milkmen, and producers, and represented as nearly as possible the milk as actually sold to the consumers. In making the averages, all samples of skimmed milk, as well as all samples standing above 17% of total solids, or under 10.75%, were deducted. The results are summarized as follows:

163

QUALITY OF MILK SOLD IN MASSACHUSETTS CITIES AND TOWNS IN WINTER AND SUMMER.

MILK.

				Dece	mber.			
	Number	•	Total Solid	s.		Pat.	304	
	of Samples.	Highest Per Cent.	Lowest Per Cent.	Average Per Cent.	Highest Per Cent.	Lowest Per Cent.	Average Per Cent.	not Fat. Average Per Cent.
Cities Towns Summary	403 99 502	16.86 15.48 16.86	10.88 12.02 10.88	13.21 13.44 13.32	8.50 6.65 8.50	2.40 3.50 2.40	4-37 4-48 4-42	8.74 8.96 8.85
				Ju	ne.			
;	Number	•	Total Solids	L		Fat.		Solids not Fat.
	of Samples.	Highest Per Cent.	Lowest Per Cent.	Average Per Cent.	Highest Per Cent.	Lowest Per Cent.	Average Per Cent.	Average Per Cent
Cities Towns Summary	311 76 387	16.90 15.71 16.90	10.75 10.99 10.75	12.67 12.63 12.65	8.80 7.10 8.80	2.10 3.00 2.10	4.03 4.09 4.06	8.54 8.54 8.54

It is interesting to note that the average for total solids of the 889 samples examined for both months stands at just 13%, of which 4.24% is fat and 8.76 is solids not fat.

Rapid Approximate Methods of Determining the Quality of Milk.—
The Lactometer.—A rough idea of the quality of milk can be gained by the use of the lactometer (page 131), but, in view of the fact that a low specific gravity may be the result either of a watered milk or of a milk high in fat, good judgment is necessary in connection with its use. A milk of good standard quality should have a specific gravity between the limits of 1.027 and 1.033. A watered milk would run below the former and a skimmed milk above the latter figure, though a milk unusually rich in fat would also run low. It should easily be apparent from the taste and appearance of the milk, whether a low specific gravity is due to watering or unusual richness in fat. The fact should also be recognized, that a milk sample may be far below the standard, and still show a specific gravity within the limits of pure milk, by skillfully subjecting the milk to both skimming and watering.

The Lactoscope.—Feser's lactoscope (Fig. 51) gives an approximation to the amount of fat in milk, and its use, especially in connection with the lactometer, is of some value. This instrument consists of a graduated glass barrel, a, into the bottom of which is accurately fitted the stopper, bearing

a white glass cylinder, having black lines thereon. Four cc. of milk are introduced into the barrel by means of a pipette, c, and water is added with thorough mixing till the translucence of the mixture is sufficient to allow the black lines to be perceptible through it. The height of the level of milk and water in the barrel a is then read off, the number indicating roughly the percentage of fat in the sample.

As in the case of the lactometer, the purity of a milk sample cannot be positively established by the lactoscope alone. For instance, a watered milk abnormally high in fat would often be found to read within the limits of pure milk, when as a matter of fact its total solids would be below standard. By a careful comparison of the readings of both the lactoscope and lactometer, however, it is rare that a skimmed or watered sample could escape detection.

Thus, if the specific gravity by the lactometer is well within the limits of pure milk, and the fat, as shown by the lactoscope, is above 3½ per cent., the sample may be safely passed as pure, or as conforming to the standard.

A normal lactometer reading in connection with an abnormally low lactoscope reading shows both watering and skimming, and with an abnormally high lactoscope reading shows a milk high in fat, or a cream. With the lactoscope reading below three, and a low lactometer reading, watering is indicated. A lactometer reading above thirty-three, and a low lactoscope reading, indicate skimming.

Heeren's Pioscope.—This instrument consists of a hard-rubber disk, having in the center a shallow receptacle, the circular rim of which is raised above the level of the disk. Into this receptacle are introduced a few drops of the milk to be tested, and a circular cover-glass containing a number of variously tinted segments is placed over the receptacle, which spreads the milk out into a thin layer, and causes it to assume a tint against the black background that can be matched with one of the colors on the glass, the various tints indicating milks of various grades from the very poorest to rich cream. This test is at best a very rough one.

Examination of the Milk Serum.—Detection of Added Water.— This may often be detected by determining the specific gravity or the degree of refraction of the milk serum, since it has been found that under fixed conditions the composition of the milk serum, or clear "whey," is more constant than that of the milk itself. Hence any considerable amount of watering is manifest from the physical constants of the serum.

In using this method the analyst should carefully work out his own

standards for comparison, by personal experiment on milk of known composition to which varying amounts of water have been added, using



FEG. 51.-Feser's Lactoscope.

the same conditions for obtaining the serum in all cases. Woodman's method * is as follows: To 100 cc. of the milk at a temperature of about 20° C. are added 2 cc. of 25% acetic acid, specific gravity 1.0350, in a

^{*} Jour. Am. Chem. Soc., 21, 1899, p. 503.

beaker, and the beaker, covered with a watch-glass, is heated in a water-bath for 20 minutes at a temperature of 70° C. After this the beaker is placed in ice water for 10 minutes and the solution filtered.

Specific Gravity.—The specific gravity of the clear filtrate, obtained by the method described above, is taken at 15° C. with the Westphal balance.

Immersion Refractometer Reading.—The instrument used is the Zeiss immersion or dipping refractometer described on pages 111 to 121. The serum, prepared as directed in a preceding paragraph, is examined in one of the small beakers accompanying the apparatus at a temperature of 20° C.

Constants of the Serum.—The three tables which follow show the variation of specific gravity and immersion refractometer reading on milk of different composition.

Analyses of whole milk submitted by the author to varying degrees of watering, up to 50% of added water, are given in the following table:

CONSTANTS	OF	MILK	AND	MILK	SERUM.	A	WHOLE	MILK
		SYSTE	MATIC	CALLY	WATERE	ο.		

	Determinations on Milk. On Milk Serum							Serum.
Added Water, Per Cent.	Total Solids, Per Cent.	Water, Per Cent.	Fat, Per Cent.	Solids not Fat, Per Cent.	Ash, Per Cent.	Specific Gravity at 15° C.	Specific Gravity at 15° C.	Immersion Refrac- tometer Reading at 20° C.
0	12.65	87.35	4.00	8.65	0.65	1.0315	1.0287	42.40
10	11.33	88.67	3.50	7.83	0.60	1.0278	1.0260	39.75
20	10.10	89.90	3.10	7.00	0.53	1.0252	1.0230	36.90
30	8.95	91.05	2.80	6.15	0.48	1.0211	1.0200	34.10
40	7.67	92.33	2.40	5.27	0.40	1.0192	1.0167	31.10
50	6.43	93.57	2.00	4.43	0.38	1.0154	1.0140	28.45

The first table on p. 167 shows a centrifugally skimmed milk, systematically watered up to 50% of added water, as in the preceding table. It will be observed that both the specific gravity and immersion refractometer readings of the serum of the whole milk, agree very closely with those of the skimmed milk in cases having a corresponding amount of added water.

The second table on p. 167 shows analyses of milk selected from a wide range of samples regularly collected and examined in the routine of food inspection by the Massachusetts State Board of Health.

MILK.

CONSTANTS OF MILK AND MILK SERUM. A SKIMMED MILK SYSTEMATICALLY WATERED.

	On Milk Serum.							
Added Water, Per Cent.	Total Solids, Per Cent.	Water, Per Cent.	Fat, Per Cent.	Solids not Fat, Per Cent.	Ash, Per Cent.	Spēcific Gravity at 15° C.	Specific Gravity at 15° C.	Immersion Refrac- tometer Reading at 20° C.
•	9.05	90.95	0.03	9.02	0.64	1.0350	1.0206	42.85
10	8.14	91.85	0.03	8.11	0.60	1.0317	1.0260	39.60
20	7.27	92.73	0.02	7.25	0.56	1.0278	1.0230	36.85
30	6.41	93.59	0.02	6.39	0.48	1.0247	1.0200	34.00
40	5.50	94.50	0.01	5.49	0.44	1.0209	1.0170	31.20
50	4.61	95 - 39	0.01	4.60	0.39	1.0172	1.0140	28.50

CONSTANTS OF MILK AND MILK SERUM. LABORATORY SAMPLES

		Determination	ons on Milk.			On Mil	On Milk Serum.		
Total Solids, Per Cent.	Water, Per Cent.	Pat, Per Cent.	Solids not Fat, Per Cent.	Ash, Per Cent.	Specific Gravity at 15° C.	Specific Gravity at 15° C.	Immersion Refractom- eter Read- ing at 20° C		
16.45	83.55	8.20	8.25		1.0255	1.0274	40.95		
15.90	84.10	7.00	8.90	0.60	1.0277	1.0285	42.00		
14.37	85.63	5.50	8.88	0.58	1.0282	1.0280	42.40		
14.17	85.83	4.85	9.32	0.62	1.0313	1.0281	44.20		
14.04	85.96	4.95	9.00	0.60	1.0303	1.0274	42.70		
13.80	86.20	5.00	8.8o	0.65	1.0302	1.0280	42.75		
13.59	86.41	4.30	9.29	0.64	1.0321	1.0285	44.50		
13.39	86.61	4.40	8.gg	0.50	1.0324	1.0285	43.70		
13.28	86.72	4.40	8.88	0.60	1.0299	1.0280	42.65		
13.12	86.88	4.00	0.12	0.59	1.0317	1.0280	43.75		
13.00	87.00	4.30	8.70	0.56	1.0310	1.0266	42.60		
12.90	87.10	3.85	9.05	o.ór	1.0318	1.0280	43.40		
12.80	87.20	4.30	8.50	0.46	1.0304	1.0277	42.70		
12.70	87.30	3.80	8.90	0.53	1.0314	1.0280	43.10		
12.63	87.37	3.50	9.13	0.65	1.0323	1.0277	43.65		
12.62	87.38	4.10	8.52	0.52	1.0298	1.0272	42.40		
12.57	87.43	3.70	8.87	0.68	1.0317	1.0278	43.45		
12.47	87.53	3.60	8.87	0.65	1.0303	1.0282	43.15		
12.36	87.64	3.20	9.16	0.55	1.0327	1.0282	43.25		
12.30	87.70	3.20	9.10	0.62	1.0327	1.0283	44.00		
12.16	87.84	4 · 35	7.8r	0.49	1.0275	1.0265	41.10		
12.00	88.00	3.40	8.60	0.62	1.0275	1.0280	41.75		
11.86	88.14	3.60	8.26	0.49	1.0306	1.0266	42.40		
11.67	88.33	3.95	7.77	0.48	1.0265	1.0240	39.30		
11.60	88.40	2.75	8.85	0.65	1.0320	1.0282	43.55		
11.50	88.50	3.45	8.05	0.51	1.0290	1.0269	41.40		
11.40	88.60	3.10	8.30	0.60	1.0297	1.0278	42.00		
11.25	88.75	2.80	8.45	0.58	1.0280	1.0274	40.90		
11.07	88.93	3.00	8.07	0.62	1.0200	1.0270	40.75		
10.69	89.31	2.95	7.74		1.0288	1.0262	39.85		
10.25	89.75	3.20	6.95	0.55	1.0230	1.0223	36.40		
8.34	91. 66	2.20	0.14	0.38	1.0224	1.0207	34.70		

A comparison of the immersion refractometer readings of the serum of milk of varying quality shows at once that the refraction of the serum is a general index to watering. A reading below 40 with the above conditions carefully observed would be suspicious of added water, though 39 might more safely be placed as a limit, below which milk could be declared fraudulently watered. The analyst need not hesitate in testifying to the presence of added water, when in addition to giving a refraction reading lower than 39 under the above conditions, the solids not fat stand below 7.3%.

The tables on page 169 are of interest, as they show, in summarized form, refractometric and analytical results from a large number of milk samples from three widely separated localities, namely, Massachusetts, New Jersey, and Great Britain:*

Nitrates.—The presence of nitrates in milk furnishes strong evidence of watering. Pure milk, free from contamination with stable filth, contains no nitrates; well water, however, often contains a sufficient amount to enable the detection of a 10% admixture in milk.

Richmond † employs for this test a solution of diphenylamin in concentrated sulphuric acid (o.1 gram to 100 cc.). One cc. of the mixture is placed in a small porcelain crucible, and a drop of the milk serum is run down the side and allowed to flow over the surface. If a blue color appears within 10 minutes, the presence of nitrates is indicated. A brownish color always forms on standing for a longer time, whether or not nitrates are present.

Patrick finds that the delicacy of the test is greatly increased by adding to the reagent a small amount of sodium chloride some time before using.

SYSTEMATIC EXAMINATION OF MILK FOR ADULTERATION.—If a large number of samples of milk have to be examined daily for adulteration, it may be an advantage to submit all to a preliminary test with the lactoscope and lactometer, excluding from further analysis, as above the standard, such samples as pass certain prescribed limits which experience has proved these tests to be capable of showing to an experienced observer, and submitting the remainder to a chemical analysis. In using such an instrument as the lactoscope for this purpose, the individual element is a most important consideration, and the use of this instrument

^{*} An. Rep. Mass. State Board of Health, 1906, p. 384.

[†] Analyst, 18, 1893, p. 272.

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Classification according to Total Solids, Per cent. Above 15 14 to 15 12 to 13 11 to 12 10 to 11	Limits. Limits. Lowest Highest Lowest Highest Lowest Highest Lowest Highest Highest Highest Highest Highest Highest Highest Highest Lowest Highest Lowest Lo	Number of Sam- Number of Sam- N spies. I Sam- Sam- Sam- Sam- Sam- Sam- Sam- Sam-	1 2 2 3 3 5 5 5 5 5 5 5 5	## 70.00 4.4.00.4.4.00	2 3 47 100 Solids not Fat, 2 2 47 17 1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2		Number of Sam-	Serior S	Laborati Per cent. Per cent. 2.9.0 % 0.0	γ	Serum at 30°05.	7 7 3 5 1 ples.	Liver 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	Cont. Cont	. a Now N 440 0 WA	O ce tammad 2 6 6 6 7 7 8 6 8 8 8 8 8 8 8 8 8 8 8 8 8
		<i>,</i>			3	23.65						-	55.53		2	3
10 to 11	Lowest	8	10.88	3.45		39.0	<u> </u>	<u> </u>	:	÷		~	10.98		8.63	%.04 %.06
<u>-</u> -	Lowest /	,	11.03	8 ;		39.9	~~	11.74	8.8	8.14	9.0		1.10		.3	39-13
~	Highest \	32	11.93	3.60	-	44.1	\	96.11	3.60	8.06	42.5	_	11.07	_	0.40	43.00
	Lowest (13 {	12.11	3-55		41.4	 	12.42	3.80	8.62	42.2	73 {	12.05	_	7.87	
_	Highest \	\	12.96	4.50	_	44.4	,			- ·	?		12.02	_	0.04	42.50
	Lowest \	-	13.43	4.35	8	43.0	~	13.16	4.10	2	7 2 7	38	12.02	_	200	2
~	Highest (1-	13 43	,	0	,	Ţ,	13.21	4.25	0.11	43.7		13.08	_	0.48	44.15
_	Towest			-	-	:	<u>~</u>	14.01	20	8.87	12.5) or (14.00	_	8.66	41.62
~	Hignest) {	14.72	5.85	9.16	43.6		14.87	_	10.25	43.60
<u>-</u>	Lowest (~	15.10	8.9	9.19	2.5		:	:	<u> </u>		-	15.12	5.90		42.63
	1				Ì	Ì	Ì.		Ì	Ì	1	Ì			Ì	
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Per cent.		nber of	ıl Solid at.	Per ce	ton el reent.	action rumat	iber of	l Solid nt.	Per ce	s, not r cent.	te tion	ber of	l Solida 1t.	Per ce	s not r cent.	noitos 18 mun
Classification according to Total Solide	Limits	Sam-	s, Per	.ta	Pat,	.D°oc.	-maS	1, Per	.31	Fat,	.D °0.	-maé	Per	ıt.	Pat,	of °C.
		Massa	chusetts S	tate Bos	urd of H		New Jan	sey State	Laborat	ory of H	ygiene.		Liverp	ool, Eng	dand.	
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		Ma	ssachusett	Massachusetts State Board of Health.	ard of Heal	Massachusetts State Board of Health. New Jer	New	Jersey Sta	te Labora	NEW Jersey State Laboratory of Hygiene.	piene.
Classification.	Limits and Average.	Number of Samples.	Total Solids. Per cent.	Fat, Per cent.	Solids not Fat, Per cent.	Solids not Refraction Number of Serum of Serum of Serum of Serum of Per cent. at 20° C. Samples.	Number of Samples.	Total Solids. Per cent.	Fat. Per cent.	Solids not Fat, Per cent.	Refraction of Serum at 20° C.
Above standard of solids (12 per cent)	Highest Lowest Average	34	14.54	3.10	10.06	45.0	48	15.37	3.00	9.20	39.6
Below standard of solids (12 per cent) but not watered.	Highest Lowest Average	31	10.27	3.55	8.7.8	39.0	33	9.12	4.00 0.35	9.90	43.3
Milk with added water	Highest Lowest	58 {	11.35	0.4.0	7-77	38.8	17 {	12.80	6.30	7.27	38.7
	* The posi	tive purity	of all sar	nples inclu	ded in thi	* The positive purity of all samples included in this table was absolutely established.	absolutely	establish	ğ.		

in the milk laboratory should be limited only to a skillful operator, accustomed to interpret its results.

The method used in the writer's laboratory has been to submit all samples to the regular test for solids, and such samples as fall below the legal standard for solids, are further examined for fat.

Total Solids, Ash, and Fat.—It is presupposed that the analyst is equipped with a sufficient number of platinum dishes for the number of milk samples daily analyzed. It is a convenience to have these dishes numbered, and instead of weighing each dish, to have a system of numbered counterweights (Fig. 52, A) corresponding to the dishes. counterweights in use by the author for this purpose are easily made from half-inch lead pipe, cut to the appropriate length and flattened, Each weight is then carefully adjusted to its appropriate dish, by trimming off the weight with a knife, or by adding bits of lead scraps, if necessary, by simply prying open in the center, inserting the required amount of scrap, and then closing by a blow of the hammer, the weight being plainly numbered before final adjustment. A rack is provided by the side of the balance-case (Fig. 52) with slits for holding the weights in their appropriate places. Such a set of counterweights is not difficult to make, requires very little care to keep in adjustment, and is an immense labor-saving device.

Details of Manipulation.—The following method of examining large numbers of milk samples is the one in use in the laboratory of the Massachusetts State Board of Health and is given in some detail, as long experience has proved it to be rapid, easy, and accurate.

From 12 to 20 samples of milk are conveniently weighed out at a sitting, the unopened sample cans or bottles being contained in a tray at the left of the operator on a low stand, another low stand and tray being at his right hand for the cans, after removing the weighed portions, and a third tray on the table at the right of the balance for the platinum dishes with the weighed samples. The analyst enters the number of the platinum dish in his note-book, or on a card,* in line with the number of the milk sample, verifies the correctness of the counterweight, and weighs out exactly 5 grams of the milk with the aid of a pipette, after first having thoroughly mixed the sample. This operation is repeated with all the samples, the platinum dishes containing the weighed amounts

^{*} Specially ruled library cards, as shown on page 172, are useful for this purpose.

of each being placed in succession on the tray, which is finally carried to the water-bath and the dishes transferred thereto. The time required for weighing out 12 samples of milk in this manner is about fifteen minutes.

The water-bath is inclosed in a hood, and the sliding front is so arranged that it can be shut down and locked, so that if the analyst has to leave

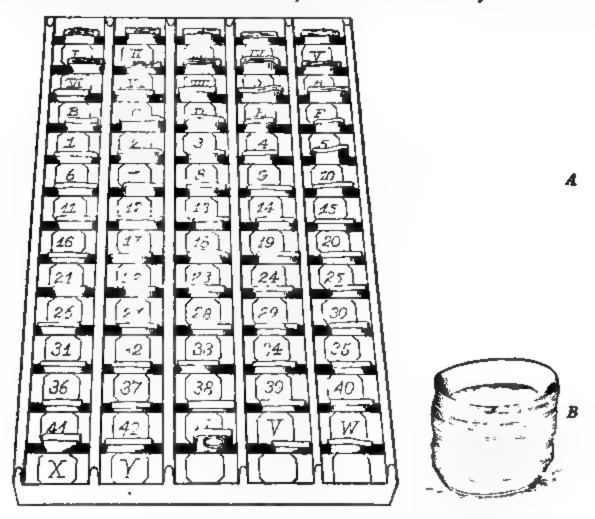


Fig. 52.—Set of Counterweights for Numbered Platinum Dishes, in a Convenient Rack.

- A. One of the Counterweights.
- B. Platinum Dishes.

the laboratory during the three hours required for the evaporation, he can swear in court that the samples could not be tampered with during his absence (see page 21).

When ready to make the second weighings for the total solids, each dish is taken from contact with the steam, and, while still hot, is wiped dry with a soft towel, till twelve of the dishes are placed on the tray, which is then taken to the balance. Experience has shown that with ordinary rapidity in weighing, twelve of the residues may be thus dealt with at a time without the need of a desiccator, the gathering of moisture during that time being inappreciable, excepting in very damp weather, when a less number of dishes should be removed at a time from the bath. In making the second weighing, and employing the counterweight as

. '							
		Date	Januar	y 6,	1904.		
Inspector's Number	No. of Dish	Wt. Residue 5 Grams.	Total Solids.	No. of Bottle	Fat.	Solids not Fat	Remarks
2642 f	1	6458	12.91				
2644	2	6530	13.06				
2646	3	6011	12.02				
2648	4	5980	11.96	3	3.25	8.71	
2650	5	7263	14.53				
2652	6	4174	8.95	4	2.50	585	Colored with
2654	7	.6823	13.65				
2656	8	. 6301	12.60				
2658	9	.6924	13.85				
2660	10	6135	12.27				
2662	11	4595	9.19	5	0.15	9.04	Marked Skanned.
2664	12	4693	9.39	6	285	6.54	
2666	13	6530	13.06				Contains Journalde hyde
2668	14	1.1452	22.90				Contains bris acid.
2670	15-	6293	12.59				
2672	16	7393	14.79				Sour
2674	17	7102	14.20				
2676	18	6010	12.02				
2678	19	4501	9.00	7	1.20	7.80	
2680	20	. 6531	13.06				
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Specimen Card for Analyst's Records of Milk Analyses. To be filed in a cabinet.

before, the exact net weight of the residue is at once ascertained and entered in the appropriate column in the note-book. Multiplied by 20 it gives at once the percentage of total solids.

It is a great saving of time to weigh out exactly 5 grams as above described. The knack of quickly measuring out the exact amount is easily acquired with practice, the 5-gram weight is the only one required for the operation with the counterweight of the dish, and the laborious figuring of percentage due to using a fraction above or below the 5 grams of milk is avoided.

Such samples as are found to stand below the standard of total solids are further examined for fat by the Babcock process (p. 136), entering the number of the fat bottle in the note-book in the appropriate column, and subsequently the percentage of fat.

Ordinarily the specific gravity is not determined, excepting in some cases of badly watered milk, when, for purposes of a check, it is customary to take the specific gravity, and calculate the solids from the gravity and the fat by Babcock's formula (p. 153), or the Richmond sliding scale, and compare the result with the figure directly determined.

The ash is rarely weighed except in special cases.

The dishes containing the dry residues are easily cleaned by first burning to an ash and cooling, after which they are treated successively with strong nitric acid, which is poured from one to another, the dishes being rinsed thoroughly with water and finally heated to redness.

A convenient device for ashing a large number of residues for purposes of cleaning the platinum dishes and for final heating is the incinerator shown in Fig. 53, made of Russia iron.

Pag. 53.—A Sheet-metal Incinerator, Specially Useful for Ashing Milk Residues.

ADDED FOREIGN INGREDIENTS.

Passing over such mythical and impossible adulterants as chalk, and the almost as rarely used substances calves' brains, starch, glycerin, sugar, etc., often discussed in manuals on milk, but with few authentic instances of their actual occurrence, the commonly found adulterants may be divided into two classes: coloring matters and preservatives.

The coloring matters almost exclusively used are annatto, azo-colors, and caramel. The preservatives commonly met with are formaldehyde, boric acid, borax, and sodium bicarbonate. Rarely salicylic and benzoic acids are found.

COLORING MATTERS.—While it is more often true that an artificially colored milk is also found to be watered, the coloring being added to cover up evidence of the watering, it is not uncommon to find added coloring matter in milk above the standard.*

About 95% of the milks found colored in Massachusetts show on analysis the fraudulent addition of water.

Statistics of the Massachusetts State Board of Health show that out of 48,000 samples of milk collected throughout the state and analyzed during nine years (from 1894 to 1902 inclusive) 342 samples or 0.7% were found to contain foreign coloring matter. Of these samples, about 67% contained annatto, approximately 30% were found with an azodye, and about 3% with caramel.

Until comparatively recently annatto was employed almost exclusively for this purpose. Caramel is least desirable of all the above colors from the point of view of the milk-dealer, in that it is difficult to imitate with it the natural color of milk, by reason of the fact that the caramel color has too much of the brown and too little of the yellow in its composition. Annatto, on the other hand, when judiciously used and with the right dilution, gives a very rich, creamy appearance to the milk, even when watered, which accounts for its popularity as a milk adulterant. Of late, however, the use of one or more of the azo-dyes has been on the increase, and so far as a close imitation of the cream color is concerned, these colors are quite as efficient as annatto.

Appearance of Artificially Colored Milk.—The natural yellow color of milk confines itself largely to the cream. An artificial color, on the contrary, is dissipated through the whole body of the milk, so that when the cream has risen in a milk thus colored, the underlying layers, instead of showing the familiar bluish tint of skimmed milk, are still distinctly tinged below the layer of the fat, especially if any considerable quantity of the color has been used. This distinctive appearance is in itself often

^{*} In one instance an azo-dye was found by the writer in a milk that contained over 17% of total solids.

sufficient to direct the attention of the analyst to an artificially colored milk, in the course of handling a large number of samples.

Nature of Annatto.—Annatto, arnatto, or annotto is a reddish-yellow coloring matter, derived from the pulp inclosing the seeds of the *Bixa* orellana, a shrub indigenous to South America and the West Indies.

A solution of the coloring matter in weak alkali is the form usually employed in milk.

Nature of "Anilin Orange."—Of the coal-tar colors employed for coloring milk, the azo-dyes are best adapted for this purpose and are most used. A few samples of these commercial "milk improvers" have fallen into the hands of the Department of Food and Drug Inspection of the Massachusetts Board of Health, and have proved, on examination, to be mixtures of two or more members of the diazo-compounds of anilin. A mixture of what is known to the trade as "Orange G" and "Fast Yellow" gives a color which is practically identical with one of these preparations, secured from a milk-dealer and formerly used by him.

For purposes of prosecution or otherwise, it is obviously best in our present knowledge of the subject to adopt a generic name such as "a coal-tar dye" or "anilin orange" * to designate this class of coloring matters in milk, rather than to particularize.

Systematic Examination of Milk for Color.—The general scheme employed by the writer for the examination of milk samples suspected of being colored is as follows: † About 150 cc. of the milk are curdled by the aid of heat and acetic acid, preferably in a porcelain casserole over a Bunsen flame. By the aid of a stirring-rod, the curd can nearly always be gathered into one mass, which is much the easiest method of separation, the whey being simply poured off. If, however, the curd is too finely divided in the whey, the separation is effected by straining through a sieve or colander. All of the annatto, or of the coal-tar dye present in the milk treated would be found in the curd, and part of the caramel. The curd, pressed free from adhering liquid, is picked apart, if necessary, and shaken with ether in a corked flask, in which it is allowed to soak for several hours, or until the fat has been extracted, and with it the annatto. If the milk is uncolored, or has been colored with annatto, on pouring off the ether the curd should be left perfectly white. If, on

^{*} The term "anilin orange" has been so commonly applied during the last eight years to any color or mixture of colors of this class in complaints in the Massachusetts courts, as to have acquired a special meaning perfectly well understood.

[†] Jour. Am. Chem. Soc., 22, 1900, p. 207.

the other hand, anilin orange or caramel has been used, after pouring off the ether the curd will be colored more or less deeply, depending on the amount of color employed. In other words, of the three colors, annatto, caramel, and anilin orange, the annatto only is extracted by ether. If caramel has been used, the curd will have a brown color at this stage; if anilin orange, the color of the curd will be a more or less bright orange.

Tests for Annatto.—The ether extract, containing the fat and the annatto, if present, is evaporated on the water-bath, the residue is made alkaline with sodium hydroxide, and poured upon a small, wet filter, which will hold back the fat, and, as the filtrate passes through, will allow the annatto, if present, to permeate the pores of the filter. On washing off the fat gently under the water-tap, all the annatto of the milk used for the test will be found to have been concentrated on the filter, giving it an orange color, tolerably permanent and varying in depth with the amount of annatto present. As a confirmatory test for annatto, stannous chloride may afterward be applied to the colored filter, producing the characteristic pink color.

Tests for Caramel.—The fat-freed curd, if colored after the ether has been poured off, is examined further for caramel or anilin orange, by placing a portion of the curd in a test-tube, and shaking vigorously with concentrated hydrochloric acid. If the color is caramel, the acid solution of the colored curd will gradually turn a deep blue on shaking, as would also the white fat-free curd of an uncolored milk, the blue coloration being formed in a very few minutes, if the fat has been thoroughly extracted from the curd; indeed, it seems to be absolutely essential for the prompt formation of the blue color in the acid solution that the curd be free from fat. Gentle heat will hasten the reaction. It should be noted that it is only when the blue coloration of the acid occurs in connection with a colored curd that caramel is to be suspected, and if much caramel be present, the coloration of the acid solution will be a brownish blue. If the above treatment indicates caramel, it would be well to confirm its presence, by testing a separate portion of the milk in the following manner.*

About a gill of the milk is curdled by adding to it as much strong alcohol. The whey is filtered off, and a small quantity of subacetate of lead is added to it. The precipitate thus produced is collected upon a small filter, which is then dried in a place free from hydrogen sulphide. A pure milk thus treated yields upon the filter-paper a residue which is

^{*} See Nineteenth Annual Report of the Mass. State Board of Health (1887), p. 183.

either wholly white, or at most of a pale straw color, while in the presence of caramel, the residue is a more or less dark-brown color, according to the amount of caramel used.

Tests for Coal-tar Dye.—If the milk has been colored with an azo-dye, the colored curd, on applying the strong hydrochloric acid in the test-tube, will immediately turn pink. If a large amount of the anilin dye has been used in the milk, the curd will sometimes show the pink coloration when hydrochloric acid is applied directly to it, before treatment with ether, but the color reaction with the fat-free curd is very delicate and unmistakable.*

Lythgoe † has shown that the amount of anilin orange ordinarily present in a milk for the purposes of coloring can be detected by adding directly to say 10 cc. of the sample an equal quantity of strong hydrochloric acid and mixing, whereupon the pink coloration is produced, if the dye is present in more than minute traces. The test is more delicate if carried out in a white porcelain dish. It had best be used as a preliminary test only, and confirmed by a subsequent test on the fat-free curd as above.

SUMMARY OF SCHEME FOR COLOR ANALYSIS.

Curdle 150 cc. milk in casserole with heat and acetic acid. Gather curd in one mass. Pour off whey, or strain, if curd is finely divided. Macerate curd with ether in corked flask. Pour off ether.

Ether Extract.

Evaporate off ether, treat residue with NaOH and pour on wetted filter. After the solution has passed through, wash off fat and dry filter, which if colored orange, indicates presence of annatto. (Confirm by SnCl₂.)

Extracted Curd.

(1) If Colorless.—Indicates presence of

no foreign color other than in ether extract.
(2) If Orange or Brownish.—Indicates presence of anilin orange or caramel. Shake curd in test-tube with concentrated hydrochloric acid.

If solution gradually turns blue, indicative of caramel. (Confirm by testing for caramel in whey of original milk.)

If orange curd immediately turns pink, indicative of anilin orange.

NATURE OF PRESERVATIVES USED IN MILK.—In most localities having pure food laws preservatives in milk are regarded as adulterants.

^{*} Occasional samples of milk colored with a coal-tar dye of a different class from those already described have recently been found in Massachusetts. In these cases the color of the separated fat-free curd does not change when treated with hydrochloric acid. The color of the curd is, however, very marked, being deep orange, bordering on the pink.

[†] Jour. Am. Chem. Soc., 22, 1900, p. 813.

Their use, however, seems to be on the decrease. Of 6,186 samples of milk examined by the Massachusetts State Board of Health during one year (1899) 71 samples, or 1.2%, were found to contain a preservative. Of these 55 were found with formaldehyde, 13 containing boric acid, borax, or a mixture of the two, and 3 contained carbonate of soda.

Comparative tests have been made in the writer's laboratory of the keeping qualities of these commonly used milk preservatives, when present in varying strength, the milk being kept during the experiment at the temperature of the room, which at that season of the year (February) was about 20° C.* The preservatives were added about five hours after milking. The samples were titrated for acidity each morning, the acidity being expressed by the number of cubic centimeters of decinormal sodium hydroxide necessary to neutralize 5 cc. of the milk.

The proportions of preservatives used in this experiment, as shown in the table on page 179, were intended to cover a wide range, from the weakest that could aid in preserving the milk up to a strength limited only by being perceptible to the taste. The table opposite shows the results.

Formaldehyde, the most commonly used preservative for milk, is sold to the trade under various names, such as "Preservaline," "Freezine," "Iceline," etc., all being dilute aqueous solutions of formaldehyde, containing from 2 to 6 per cent of the gas, being nearly always diluted from the 40% solution known as formalin. These preparations are usually accompanied by directions, which specify the amount to be used, varying from a tablespoonful of the solution in 5 to 10 gallons of the milk. It is commonly used in the strength of 1 part of the gas in 20,000, and rarely less than 1 part in 50,000. The antiseptic power of formaldehyde increases in a marked degree as the strength of the preservative is increased. Milk treated with 1 part in 10,000, for instance, according to the table was found to keep sweet $5\frac{1}{2}$ days. In the strength of 1 part to 5000, the milk did not curdle for 101 days, while 1 part of formaldehyde to 2500 parts of milk kept the milk from curdling for 55 days, the acidity up to that time being nearly normal.

Formaldehyde is thus shown to be decidedly the most efficient of all milk preservatives, besides being inexpensive and convenient to use.

Whether the growth of other bacteria than those that produce lactic fermentation is inhibited by formaldehyde in milk is not definitely settled. The claim has been made that pathogenic varieties are destroyed by its use.

^{*} Thirty-first Annual Report Mass. State Board of Health, 1899, p. 611.

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Whether or not formaldehyde in milk is harmful to processes of digestion, when present in the amount commonly used, is still an open question.*

Carbonate and Bicarbonate of Soda.—These substances are occasionally used in milk, though, as the above table shows, they possess little or no value as milk preservatives. They do, however, serve to neutralize the acidity of slightly soured milk and to postpone the time of actual curdling.

Salicylic and Benzoic Acids, in view of the much more efficient antiseptics at hand, are now rarely used as milk preservatives, though the analyst should be on the outlook for them. Salicylic acid is a poor milk preservative, in view of the fact that it affects the taste of the milk, when present in sufficient quantity, to be of service.

Detection of Formaldehyde.—Hydrochloric Acid Test.†—Commercial hydrochloric acid (specific gravity 1.2) containing 2 cc. of 10% ferric chloride per liter is used as a reagent. Add 10 cc. of the acid reagent to an equal volume of milk in a porcelain casserole, and heat slowly over the free flame nearly to boiling, holding the casserole by the handle, and giving it a rotary motion while heating to break up the curd. The presence of formaldehyde is indicated by a violet coloration, varying in depth with the amount present. In the absence of formaldehyde, the solution slowly turns brown. By this test I part of formaldehyde in 250,000 parts of milk is readily detected before the milk sours. After souring, the limit of delicacy proves to be about 1 part in 50,000.

Various aldehydes, when introduced into milk, give color reactions under the above treatment, but formaldehyde alone gives the violet coloration, which is perfectly distinguishable and unmistakable.

Hehner's Sulphuric Acid Test.—To 5 to 10 cc. of milk in a wide testtube add about half the volume of concentrated commercial sulphuric acid,‡ pouring the acid carefully down the side of the tube, so that it forms a layer at the bottom without mixing with the milk. A violet zone at

preservative, a weak solukes the following remark-

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^{*} Milk-dealers are led to believe, by artful dealers in preservative preparations, that the chemist cannot detect them. The manufi tion of formaldehyde, issues an attractive ; able claims:

[&]quot;It is not an adulterant. It immediat as soon as it has rendered all the bacteris ence in milk, quantitatively or otherwise.'

[†] Annual Report Mass. State Board of so 1899, p. 699.

[‡] The coloration produced seems to depend on the presence of iron salts in the acid, hence the use of commercial acid is recommended. If only pure acid is available, a little ferric chloride should be added

the junction of the two liquids indicates formaldehyde. This test may be combined with the Babcock test for fat, noting whether a violet color forms on addition of the commercial sulphuric acid to the milk in the test bottle.

Confirmatory Tests with Distilled Milk.—If it is desired to confirm the above tests by further evidence, 100 to 200 cc. of the milk sample are subjected to distillation, and the first 20 cc. of the distillate are used for testing.

- (1) To a few drops of this distillate in a test-tube add a drop of Schiff's reagent.* In presence of any aldehyde, a pink coloration will soon be perceptible, deepening in intensity on standing.
- , (2) Add to 5 cc. of the milk distillate a few drops of a 1% aqueous solution of resorcin or phenol, and proceed as directed on page 820 (preservatives). The crimson color indicates formaldehyde, and not other aldehydes.
- (3) Use 1 or 2 cc. of the milk distillate and apply the phenylhydrazine test, page 820.
- (4) A small amount of the distillate from milk (which prior to distilling is acidified slightly with sulphuric acid to fix any free ammonia) is treated with a few drops of Nessler's reagent.† Traces of formaldehyde produce a yellow coloration, while if considerable formaldehyde be present, the color darkens on standing and a grayish precipitate may be formed.

Determination of Formaldehyde in Milk.‡—To 100 cc. of milk add 1 cc. of 1:3 sulphuric acid and subject to distillation in a 500-cc. Kjeldahl nitrogen-flask, using a low circular evaporating burner to avoid frothing. According to Smith, the first 20 cc. of the distillate, or one-fifth the original volume, contain very nearly one-third of the total formaldehyde. Collect 20 cc. of the distillate and determine the formaldehyde therein by the potassium cyanide method, as follows: §

Treat 10 cc. of tenth-normal silver nitrate with 6 drops of 50% nitric acid in a 50-cc. flask, add 10 cc. of a solution of potassium cyanide containing 3.1 grams of KCN in 500 cc. of water, and make up to the 50-cc. mark. Shake, filter, and titrate 25 cc. of the filtrate with tenth-normal ammonium sulphocyanate, using ferric chloride as an indicator.

^{*} Table of reagents, No. 226.

[†] Table of reagents, No. 187.

[‡] Smith, Jour. Am. Chem. Soc., 25, 1903, pp. 1032 and 1037.

[§] Zeits. anal. Chem., 36, pp. 18-24.

Theoretically 7.6 grams per liter. On account of the deliquescent nature of the salt weigh out 8 grams, make up to a liter, and titrate against tenth-normal silver nitrate for its exact value, using ferric chloride as an indicator. Sutton, Volumetric Analysis, 8th Ed. p. 155.

Acidify another portion of 10 cc. of tenth-normal silver nitrate with nitric acid, add 10 cc. of the potassium cyanide solution to which the above 20 cc. of the formaldehyde distillate has been added. Make up the whole to 50 cc., filter and titrate as before 25 cc. of the filtrate with tenth-normal ammonium sulphocyanate for the excess of silver.

The amount of potassium cyanide used up by the formaldehyde, in terms of tenth-normal ammonium sulphocyanate, is found by multiplying by two the difference between the two results, and the total formal-dehyde is calculated by multiplying by 3 the amount found in the 20 cc. of distillate.

The reaction that takes place between the formaldehyde and the potassium cyanide probably results in the formation of an addition product as follows:

$CH_2O + KCN = KO.CH_2CN.$

Detection of Boric Acid.—This is best accomplished by the turmeric-paper test applied either directly to the milk or to the ash (page 823). In the former case 10 cc. of milk are thoroughly mixed with 6 drops of concentrated hydrochloric acid, after which the tumeric paper is moistened with the mixture and dried.

Determination of Boric Acid.—Use the method of Thompson.* Add 10 cc. of a 1:1 solution of sodium hydroxide to 100 cc. of the milk, evaporate to dryness in a platinum dish, and proceed as described on page 823.

Detection of Carbonate and Bicarbonate of Soda. — The addition of carbonates is manifest by the effervescence caused by treating the milk-ash with acid. Effervescence in the milk-ash is quite perceptible, when as much as 0.05% of sodium carbonate is present.

Schmidt's method of detecting sodium carbonate or bicarbonate, when present to the extent of 0.1% or more, is as follows: Ten cc. of milk are mixed with an equal volume of alcohol, and a few drops of a 1% solution of rosolic acid are added. If carbonate is present, a rosered color will be produced, while pure milk shows a brownish-yellow coloration. The suspected sample thus treated should be compared with a similarly treated sample of pure milk at the same time.

Detection of Benzoic Acid.—Shake 5 cc. of hydrochloric acid with 50 cc. of the milk in a flask. Then add 150 cc. of ether, cork the flask and shake well. Break up the emulsion which forms by the aid of a centrifuge, or, in the absence of a centrifuge, extract the curdled milk by gently shaking with successive portions of ether, avoiding the forma-

^{*} Jour. Soc. Chem. Ind., 12, p. 432.

tion of an emulsion. A volume of ether largely in excess over that of the curdled milk has been found to be less apt to emulsionize.* Transfer the ether extract to a separatory funnel, and separate the benzoic acid from the fat by shaking out with dilute ammonia, which takes out the former as ammonium benzoate. Evaporate the ammonia solution in a dish over the water-bath till all free ammonia has disappeared, but before getting to dryness, add a few drops of ferric chloride reagent.

The characteristic flesh-colored precipitate indicates benzoic acid. Care should be taken not to add the ferric chloride till all the ammonia has been driven off, otherwise a precipitate of ferric hydrate is formed.

Detection of Salicylic Acid.—(1) To 50 cc. of the milk add 1 cc. of acid nitrate of mercury reagent (p. 147), shake and filter. The filtrate, which should be perfectly clear, is then shaken with ether in a separatory funnel, the ether extract evaporated to dryness, and a drop of ferric chloride reagent applied. If salicylic acid be present, a violet color will be produced. In carrying out the test it should be noted that a small portion only of the salicylic acid is in the filtered whey, the larger part being left in the curd. The color test is, however, so delicate as to show its presence, when an appreciable amount is used.

(2) Proceed exactly as directed for benzoic acid (p. 182). On applying the ferric chloride to the final solution, after evaporation of the ammonia, a violet color shows the presence of salicylic acid.

Routine Inspection of Milk for Preservatives.—It was the writer's custom in Massachusetts to examine all the samples of milk collected during the months of June, July, August, and September for the commonly used preservatives, in addition to the regular analysis for total solids and fat. The number of samples thus examined amounted to upwards of 500 per month, varying from 10 to 60 per day. The results of such an examination during four years are thus shown: †

		ILLUIN	CVILLIA	J 111 14111	, , , , , , , , , , , , , , , , , , ,		
Year.	Samples Examined.	Number containing Form- aldehyde.	Per Cent containing Form- aldehyde.	Number containing Boric Acid.	Per Cent containing Boric Acids	Number containing Carbonate.	Total containing Preserva- tive.
1898	1046	26	2.5 2.6	11	1.0	4	41
1899	2105 2018	55 61	3.0	13 6	0.6 0'3	3	71 67
1901		42 29	1.9 1.5	12 14	0.5 0.7	_	54 43
Totals	9257	213	2.3	56	0.6	7	376

PRESERVATIVES IN MILK.

^{*} When this process is used the ether may readily be recovered by distillation.

[†] An. Rep. Mass. State Board of Health, 1902, p. 474; Analyst's Reprint, p. 22.

Such a system by no means involves a large amount of time or labor, and is really essential before passing judgment upon the purity of the milk, since, unlike added color, there is nothing in the physical appearance of the milk to suggest the presence of preservatives, nor are they rendered apparent by the taste, if skilfully used.

The methods employed are carried out as follows:*

- (1) Formaldehyde.—After having been examined for total solids and fat, the milk samples are arranged in order in their original containers, and about 10 cc. of each sample are poured into a casserole and tested in succession by means of the hydrochloric acid and ferric chloride test (p. 180). A large stock bottle, which may be fitted with a siphon if desired, is kept on hand containing the hydrochloric acid reagent. Less than one minute is required in making the formaldehyde test for each sample.
- 2. Carbonate and Boric Acid.—These tests have been so simplified as to be, as it were, a side issue in the process of cleaning the platinum dishes used for the determination of total solids. The various residues from the total solids are burnt to an ash in the original numbered dishes in succession, these dishes, after incineration, being arranged side by side on a flat tray. By means of a pipette, one or two drops of dilute hydrochloric acid are introduced into each dish in succession, noting at the time any effervescence that may ensue, which is in itself an indication of sodium carbonate. After every milk ash has been acidulated, a few cubic centimeters of water are added to each dish by means of a washbottle, the dissolving of the ash being hastened by giving a rotary motion to the tray containing the dishes. A strip of turmeric-paper is then allowed to soak for a minute or so in each dish, after which it is withdrawn from contact with the solution and allowed to adhere to the side of the dish above the liquid, where it remains until dry. If the paper when dry is of a deep cherry-red color, turning a dark olive when treated with dilute alkali, the presence of boric acid is assured. These methods are, of course, preliminary tests for quickly singling out the preserved samples. Such confirmatory tests as are desired may in all cases be employed.

Another method of drying the strips outside the dishes is as follows: In a part of the laboratory free from dust, two long sections of glass rod or tubing are placed in parallel lines over a strip of filter-paper,

^{*} Leach, Analyst, XXVI, p. 289. An. Rep. Mass. State Board of Health, 1901, p. 447 Food and Drug Reprint, p. 27.

with numbers marked on the paper at close intervals corresponding to the numbers of the platinum dishes. The strips of turmeric-paper, after soaking, are removed from the dishes and placed across the glass tubes, over the numbers corresponding to those of the dishes from which they were taken. Here they are allowed to stand till dry, being kept in position by a third section of tube or rod placed over them. When dry, the color of the turmeric strips will indicate whether or not boric acid is present, and also the position will show in what sample to look for it.

Cane Sugar.—This is alleged to be added for the purpose of increasing the total solids of milk, but if present to any marked degree, it could hardly fail of detection by reason of the sweet taste imparted to the milk. Cane sugar in milk may be detected * by boiling 5 to 10 cc. of the sample with about 0.1 gram of resorcin and a few drops of hydrochloric acid for a few minutes. In the presence of cane sugar, a rose-red color is produced.

According to Richmond, cane sugar may be estimated by first ascertaining the total polarization of the sample as in the estimation of milk sugar (p. 147). The milk sugar is then determined by Fehling's solution (pp. 149 to 150) either volumetrically or gravimetrically. The difference between the anhydrous milk sugar found by the latter, or Fehling method, and that calculated by dividing the polarization by 1.217 will give the percentage of cane sugar present.

Cotton's † method of detecting cane sugar, when present to the extent of 0.1%, consists in mixing in a test-tube 10 cc. of the suspected milk with 0.5 gram of powdered ammonium molybdate, and adding to the mixture 10 cc. of dilute hydrochloric acid (1 to 10). Ten cc. of milk of known purity, or 10 cc. of a 6% solution of milk sugar are similarly treated by way of comparison. Both tubes are placed in a water-bath and the temperature gradually raised to 80° C. If cane sugar is present, an intense blue coloration is produced, while the genuine milk or the solution of milk sugar remains unchanged at the temperature of 80°. If the temperature is raised to the boiling-point, however, the pure milk or milk sugar solution may also turn blue.

Detection of Starch in Milk.—A small quantity of milk is heated in a test-tube to boiling, cooled, and a drop of iodine in potassium iodide added. A blue coloration indicates starch.

^{*} Richards and Woodman, Air, Water, and Food, p. 166.

[†] Abs. Analyst, 1898, p. 37.

Condensed Skimmed Milk as an Adulterant.—The use of condensed unsweetened skimmed milk to raise the solids of a skimmed or watered milk above the standard has been noted in Massachusetts. This sophistication is rendered apparent by the abnormally high solids not fat of the sample, which in some instances have exceeded 11%. A solid not fat in excess of 10% is suspicious of this form of adulteration. By fixing a legal standard for both fat and solids not fat, such tampering with milk may readily be checked.

Analysis of Sour Milk.—It occasionally becomes necessary for the analyst to deal with samples of sour milk, especially in the summer-time, when the milk has been brought from a long distance. While the process of lactic fermentation results in the formation of traces of volatile acids, unless the sample has become so badly curdled as to render an even homogeneous mixture of the various parts impossible, a fair determination of the solids and fat can readily be made. Experience has proved that, excepting in instances of milk so badly soured as to have become actually putrid, the analysis of sour milk, if carefully made, should not differ materially from that of the same milk before souring.

Care must be taken to secure an even emulsion of the curd and whey. This may sometimes be accomplished by repeatedly pouring the sample back and forth from one container to another. Again, it is sometimes necessary to use an egg-beater of the spiral wire pattern, which preferably should easily fit the can or milk-container. Unless a fine, even emulsion can be secured, it is impossible to make a satisfactory analysis of sour milk. With such an emulsion results can be relied on.

In measuring portions of the thoroughly mixed sample of sour milk for analysis, a pipette should be used having a large opening.*

CONDENSED MILK.

Canned condensed milk has become a very important article of food, its use having increased considerably during the last few years. The universally accepted meaning of the term "condensed milk" in this country is milk both condensed and preserved with cane sugar, being what is commonly known in England as "preserved milk." The unsweetened variety is more often termed "evaporated cream" and sold as such. It is, however,

^{*} A pipette open to the full size of the tube is convenient for this work.

as found on the market usually nothing better than condensed ordinary milk, having no added sugar, and has generally no resemblance in composition to cream other than in consistency.

Condensed milk is usually prepared by boiling milk in vacuum-pans under diminished pressure to the proper degree of concentration. Upwards of 350 samples of sweetened condensed milk have been analyzed in full in the laboratory of the Massachusetts State Board of Health in the course of eight years, representing no less than 110 brands, together with about 30 samples (representing 8 brands) of the unsweetened variety.

In view of the fact that a considerable number of the condensed-milk samples are shown by their analysis to have been produced from skimmed milk, the fat content in the samples analyzed varying from a mere trace to 12%, it is obvious that the typical composition of condensed milk could not fairly be shown by giving maximum, minimum, and mean results from the entire tabulated series, nor would it be possible to draw a hard-and-fast line excluding certain samples known to be adulterated in making up the averages. It has therefore been thought best to select a few typical brands and give their analyses in full.

COMPOSITION OF SWEETENED CONDENSED MILK.

Points to be Noted.	Total Solids, Per Cent.	Water Per Cent.	Milk Solids, Per Cent.	Cane Sugar, Per Cent.	Milk Sugar, Per Cent.	Pro- teins, Per Cent.	Fat, Per Cent.	Ash, Per Cent.	Fat in Origi- nal Milk, Per Cent.
High in fat, much added		0.		0			1		
sugar	79.17	20.83	31.32	47.05	9.57	7.95	12.00	1.80	4.00
Low fat, high milk sugar		32.30	30.27	30.43	0.30	10.70	11.40	1.73	5.03
low proteins		30.70	31.83	37.47	16.75	6.34	7.20	1.54	2.77
Normal constituents		3	3-1-3	3,14,	***,*		,,,,,		
		25.71	32.37	41.92	11.97	8.46	10.65	1.20	4.56
Condensed from skimmed						1		•	' '
milk	69.30	30.70	29.15	40.15	11.89	12.15	3.06	2.05	1.11
Condensed from centrifu-									l_
gally skimmed milk	69.06	30.94	25.88	43.18	11.55	111.78	0.00	2.46	Trace

COMPOSITION OF UNSWEETENED CONDENSED MILK.

Points to be Noted.	Total Solids, Per Cent.	Water, Per Cent.	Milk Sugar, Per Cent.	Pro- teins, Per Cent.	Fat, Per Cent.	Ash, Per Cent.	Fat in Original Milk, Per Cent.	No. of Times Con- densed.
High in fat. Low in proteins. Normal constituents throughout* Condensed from skimmed milk	36.00	64.co	10.65	11.63	14.00	1.72	4.61	2.6
	31.25	86.75	13.40	7.02	9.60	1.23	4.18	2.3
	28.16	6g.24	9.85	8.66	8.10	1.55	3.68	2.2
	35.17	64.83	13.90	15.37	4.20	1.70	1.28	3.3

^{*} Can be taken as being very near the average for all constituents in honest condensed milk of fair quality.

In the case of sweetened condensed milk it will be observed that the proteins as a rule run considerably lower than the sugar, whereas in ordinary cow's milk the percentage of proteins and milk sugar are more nearly alike. In making the above analyses all the reducing sugar was reckoned as milk sugar, whereas it is possible that a small amount of the cane sugar is inverted in the process of manufacture, and thus increases the amount of reducing sugar.

U. S. Standards.*—Standard condensed milk and standard sweetened condensed milk are condensed milk and sweetened condensed milk respectively, containing not less than 28% of milk solids, of which not less than 27.5% is milk fat. Standard condensed skim-milk is skimmilk from which a considerable portion of water has been evaporated.

ANALYSIS OF CONDENSED MILK.

Preparation of the Sample.—For the analysis of condensed milk the following system of procedure has been adopted in the laboratory of the Massachusetts State Board of Health. The sample is first thoroughly mixed, best by transferring the entire contents of the can to a large evaporating-dish, and working it thoroughly with a pestle till homogeneous throughout. Forty grams of the mixed sample are weighed out, preferably in a tared weighing-tray for sugar analysis, transferred by washing to a graduated 100-cc. sugar-flask (or if desired it is weighed directly into the flask) and made up to the mark with water.

Total Solids.—An aliquot part of this mixed solution is further diluted with an equal amount of water, and 5 cc. of the diluted mixture, corresponding to 1 gram of the condensed milk, is pipetted into a tared platinum dish, such as is used for ordinary milk, the pipette being rinsed into the dish by means of a wash-bottle. The dish with its contents is then placed on the water-bath, and distilled water added by the wash-bottle till the dish is nearly full. It is allowed to remain in contact with the live steam of the water-bath for at least two hours after the last traces of water have been evaporated off to leave an apparently dry residue. It is then transferred to a desiccator, cooled, and weighed.

It is of great importance to have the sample very dilute to properly determine the total solids in this manner. Formerly the sample was evenly distributed over asbestos fiber in the dish, but more accurate results were found possible by the above method. The character of the residue should be noted. It should not, excepting in the case of a

^{*} U. S. Dept. of Agric., Off. of Sec., Circ. 19.

skimmed milk, be caked down hard and glossy on the bottom of the dish, but, if the operation is properly carried out, should have a well-separated fat layer at the top, and the residue should resemble in appearance that from ordinary milk. This result is accomplished by the extreme dilution of the sample.

Ash.—The residue from the total solids as above obtained is carefully burnt, cooled, and weighed as in the case of ordinary milk (p. 134).

When the total solids are not to be determined, as in cases where the quality of the milk used in preparation of the sample is decided by the fat and ash alone (see p. 192), 12.5 cc. of the above 40% solution, corresponding to 5 grams of the sample, are evaporated to dryness on the water-bath, and the residue burnt to an ash in the muffle or over a low flame.

Fat.—The Author's Method.*—Fifteen cc. of the 40% solution prepared as above described, corresponding to 6 grams of the original condensed milk, are measured into an ordinary test-bottle of the Babcock centrifuge. This is filled nearly to the neck with water, and 4 cc. of a solution of copper sulphate of the strength of Fehling's copper solution are added. The contents are thoroughly shaken, and the precipitated proteins, carrying with them the fat are rapidly separated out by whirling the fat bottle in the centrifuge, preferably without heating. The writer prefers an electric centrifuge of the Robinson type (p. 137) for this purpose, as the heat of the steam-driven machine cakes the precipitate down, so that it is harder to wash. If desired, the precipitate may be allowed to settle out of itself, which it does more quickly in the cold.

The supernatant liquid containing the sugar is drawn off by means of a pipette of large capacity, having a stem sufficiently small to pass easily into the neck of the milk-bottle, a small wisp of absorbent cotton being first twisted over the bottom of the pipette to serve as a filter. On withdrawing the pipette with the sugar solution, the cotton is wiped off into the bottle by rubbing against the inner side.

The precipitated proteins and fat are given two additional washings, as above, by shaking thoroughly with water introduced nearly to the neck of the bottle, separating out in each case by centrifuge or by settling, and finally removing the washings with the pipette, two of such extra washings being found nearly always sufficient to remove all the sugar. If the precipitate is caked down hard after treatment with the centrifuge,

^{* 28}th An. Rep. Mass. State Board of Health, 1896, p 630, and Jour. Am. Chem. Soc., 22, 1900, p. 589.

it may be necessary to employ a stiff platinum wire as a stirrer to aid in mixing with the wash-water.

Finally, enough water is added to amount approximately to the normal volume of 17.6 cc. usually employed for the Babcock test, 17.5 cc. of sulphuric acid are added, and the test continued from this point on as in the ordinary Babcock process of milk-testing, multiplying the reading obtained by three to give the correct percentage of fat in the sample.

For condensed milk containing no added cane sugar, these precautions are, of course, unnecessary, the ordinary Babcock method being directly employed with a weighed portion of the milk.

Proteins.—Five cc. of the 40% solution originally prepared, corresponding to 2 grams of the condensed milk, are diluted further to about 40 cc., and just enough of the Fehling copper solution is added, drop by drop, to precipitate the albuminoids, taking care to avoid a large excess. As a rule, 0.6 cc. of copper solution is ample for this. Nearly neutralize with sodium hydroxide, stopping just short of alkalinity, i.e., leaving the solution still slightly acid. An excess of alkali tends to dissolve the casein and cause turbidity in the filtrate. Pass through a weighed filter-paper, wash, dry in an air-oven at 100° C., and weigh. The filter with the dry precipitate is then carefully burnt in a porcelain crucible, and the difference between the weight of the dry precipitate and the weight of the ash is the weight of the proteins and fat. Expressing this in percentage, and deducting from it the per cent of fat previously obtained, the result is the per cent of proteins.

Milk Sugar.—Volumetric Process.—The filtrate and the washings from the preceding operation are made up to 100 cc. with water, and the amount of reducing sugar, obtained volumetrically by Fehling's solution, is reckoned as milk sugar. The titration is conducted in the manner described on p. 591.

Assuming the solution to be exactly of the strength above described, the milk sugar is calculated as follows: $\frac{100\times0.067}{S\times0.02} = L$, where L is the per cent of lactose or milk sugar, and S the number of cc. of milk solution, prepared as above required to reduce 10 cc. of Fehling's solution. Calculation may be avoided by the use of the following table, which may be employed when the above details are minutely rarried out:

PER CENT MILK SUGAR CORRESPONDING TO NUMBER OF CUBIC CENTIMETERS USED.

Strength	a.f	coliftian	•	are me	in		•
Strength	Οī	solution	2	grains	111	100 (æ.

Cu. Cm.	Per Cent.						
18.0	18.61	25.0	13.40	32.0	10.47	39.0	8.59
18.5	18.10	25-5	13.14	32.5	10.31	39-5	8.49
19.0	17.63	26.0	12.89	33.0	10.15	40.0	8.37
19.5	17.18	26.5	12.64	33-5	10.00	40.5	8.27
20.0	16.75	27.0	12.41	34.0	9.85	41.0	8.17
20.5	16.34	27-5	12.18	34-5	9.71	41.5	8.07
21.0	15.95	28.0	11.97	35.0	9-57	42.0	7.98
21.5	15.58	28.5	11.75	35-5	9-43	42.5	7.88
22.0	15.22	29.0	11.55	36.0	9.30	43.0	7.78
22.5	14.89	29.5	11.35	36.5	9-17	43-5	7.70
23.0	14.56	30.0	11.16	37.0	9.05	44.0	7.61
23.5	14.25	30.5	10.89	37-5	8.93	44-5	7-53
24.0	13.95	31.0	10.80	38.0	8.8r	1	1
24.5	13.67	31.5	10.63	38.5	8.70	1	1

Gravimetric Methods.—Lactose may be determined in the 40% solution of the condensed milk by the O'Sullivan-Defren method (page 150), the Soxhlet method (page 150), or the Munson and Walker method (page 151), the solution being treated exactly as if it were milk.

Cane Sugar.—This is obtained by difference, deducting the milk solids (the sum of the milk sugar, proteins, fat, and ash) from the total solids first obtained.

Fat in Sweetened Condensed Milk.—Judgment as to the quality of a given brand of condensed milk is naturally based more on its fat content than on any other one factor, in that, of all its constituents, the fat is the only one that can conveniently be tampered with to the detriment of its value as a food. Hence, an accurate method for the determination of the most important ingredient, the fat, is of great importance.

The Babcock process without modification cannot be used, on account of the charring by the sulphuric acid acting on the cane sugar.

The Adams-Soxhlet method is unreliable, because the large amount of cane sugar is again a disturbing factor, enclosing the fat particles so firmly, when dried on the extraction coil, as to render its complete removal by the extracting ether difficult if not impossible.* In 1895 the writer's method described on page 189 was devised, and with certain minor modifications has been used ever since with highly satisfactory results, proving itself to be not only much quicker than the Adams-Soxhlet extraction

^{*} When ordinary ether is used for the Soxhlet extraction, the results may not appear too low because the alcohol and water present in the ether dissolve not only fat but also sugar, which goes in with and is weighed as fat. With ether carefully dehydrated and freed from alcohol or with benzine or petroleum ether, the fat results will always be found far too low when the extraction is conducted under ordinary conditions.

method and easier of manipulation, but, indeed, more accurate, by reason of the fact that the cane sugar with all its attendant troubles is first eliminated.

Calculation of Fat in Original Milk.—The "fat in the original milk," as expressed in the tables on page 187, was calculated by assuming a percentage of solids not fat of 9.3 in the original milk, this being the standard fixed by the Massachusetts law. Calculate first the fat and the milk solids to the basis of the cane-sugar-free sample. This is done by dividing the per cent of each as found in the sample by 100 less the percentage of cane sugar, and multiplying the result by 100. Ascertain the difference between the milk solids and the fat thus obtained in the cane-sugar-free sample, and divide this percentage of milk solids not fat by 9.3. The result is the "number of times condensed" (if cane sugar were not present as a diluent).

The per cent of fat in the cane-sugar-free sample, divided by the number of times condensed, as above obtained, gives the percentage of fat in the original milk.

The above calculation from the solids not fat of the factor designated as "the number of times condensed," necessitates determinations of fat, ash, proteins, and milk sugar, in fact, a complete analysis of the sample.

A simpler method of calculating the "number of times condensed," involving determinations of fat and ash only in the sample, consists in dividing the per cent of ash found in the condensed milk by 0.7, this figure being the assumed ash of normal, standard milk. Then, by dividing the fat in the sample by the "number of times condensed" as last calculated, the result is the fat in the original milk. If this is found to be well below 3%, there is reason to suspect that skimmed milk was used in its preparation.

The "fat in the original milk" as thus calculated is, of course, an arbitrary factor and is useful only in deciding whether or not skimmed milk has been used in preparing the sample. By assuming the above very reasonable figures for the solids not fat, or for the ash of natural milk (according to which method is used for calculation), it is readily seen that the highest result is obtained for the "fat in the original milk" and hence the benefit of the doubt as to the use of skimmed milk is given to the manufacturer.

Other Methods for Protein and Cane Sugar.—If desired, the proteins of condensed milk can be calculated from the total nitrogen obtained by the Gunning or Kjeldahl method, as in ordinary milk (p. 145).

Bigelow and McElroy's Polarimetric Method for Cane Sugar.*—26.048

^{*} Jour. Am. Chem. Soc., 15, p. 668.

MILK. 193

grams of the mixed sample of condensed milk are transferred to a 100-cc. graduated sugar-flask and dissolved in water, which is boiled to make sure of normal rotation. The solution is then clarified by the addition of an acetic acid solution of mercuric iodide * and, if necessary, alumina cream, the volume is made up to 100 cc., shaken, and filtered through a dry filter. Rejecting the first part of the filtrate, a further portion is polarized. For inversion, another sample of 26.048 grams is weighed out as before and dissolved, but before clarifying, is heated to 55° C. and treated with half a cake of compressed yeast, the heating with the yeast being continued at 55° for five hours. The clarifying solution is added before cooling, and, after cooling, making up to 100 cc., and filtering as before, the invert reading is obtained with the polariscope. By this process of yeast inversion the cane sugar only is inverted, the lactose remaining unchanged. It is best to work with several samples and use the mean of the readings both for direct and invert figures. It is also best to use the double dilution method (p. 140) to compensate for the volume of the precipitated fat and proteins.

The per cent of cane sugar is calculated by the formula of Clerget,

$$S = \frac{a - b}{142.66 - \frac{t}{2}},$$

S being the per cent of cane sugar,

a the direct reading,

b the invert reading and t the temperature at which the observation is made.

The above process presupposes the absence of invert sugar in the sample, a supposition which Wiley claims it is fair as a rule to assume.

CREAM.

Composition.—Cream varies in composition according to the method by which it is obtained, i.e., whether (1) by allowing it to separate from the milk set in shallow pans, whence it is removed by hand-skimming, (2) by setting in deep vessels surrounded by cold water (as for example in the "Cooley" creamer) the skimmed milk being commonly drawn off from below, or (3) by the centrifugal separator. Most of the heavy cream found in the market at the present time is the product of the third or separator process.

^{*} Prepared by dissolving 53 grams of potassium iodide, 22 grams mercuric chloride, and 32 cc. of strong acetic acid in water and making up to 1 liter.

Number of Analyses	Authority		Water.	Proteins.	Pat.	Sugar.	Ash.	Total Solids.	Solids not Pet.
	König	Mean	68.82	3.76	22.66	4-23	0.53	31.18	8.42
18	Leach								
18		Mean Maximum Minimum	51.68 83.29 70.50		42.02 21.60 8.60			48.32 29.50 16.71	6.30 9.30 7.22
	exigue 4 6 8	asirany 46 König 18 Leach	46 König Mean 18 Leach Maximum Minimum Mean 18 Leach Maximum Minimum Minimum	46 König Mean 68.82 18 Leach Maximum 54.80 Minimum 46.76 Mean 51.68 Maximum 83.29 Minimum 70.50	46 König Mean 68.82 3.76 18 Leach Maximum 54.80 Mean 51.68 Maximum 83.29 Minimum 70.50	46 König Mean 68.82 3.76 22.66 18 Leach Maximum 54.80 46.40 Minimum 46.76 38.10 Mean 51.68 42.02 Maximum 83.29 21.60 Minimum 70.50 8.60	18 Leach Maximum 54.80 46.40 Mean Maximum 46.76 38.10 Mean Maximum 83.29 21.60 Minimum 70.50 8.60	## ## ## ## ## ## ## ## ## ## ## ## ##	18 Leach Maximum 54.80 46.40 53.24 Minimum 46.76 38.10 42.02 48.32 Minimum 70.50 8.60 16.71

COMPOSITION OF CREAM.

Methods of Analysis.—The total solids, ash, sugar, and proteins are determined by similar processes to those used in milk analysis.

Fat.—The most convenient method of estimating fat is slightly modified from the regular Babcock process. The specific gravity of cream



Fig. 54.—Varieties of Babcock Test Bottle for Cream.

varies between such wide limits that it is best to weigh rather than measure the sample. Two varieties of cream bottle are in common use (Fig. 54) for the Babcock process, with a capacity for measuring 25 to 30 per cent of fat.

Approximately 10 grams of the well-mixed cream sample are transferred to one of these cream bottles, previously tared, and the weight of the cream accurately obtained. A convenient pipette to use for the purpose is one the end of which has been broken off to the full size of the tube.

Fig. 55 shows a cream-test scale specially designed for weighing the sample, provided with a sliding poise for counterbalancing the bottle, and a second weight for weighing the cream. The scale is delicate to 0.01 gram when loaded.

Five to 6 cc. of water are added to the cream in the bottle, after which the regular amount of sulphuric acid used in the Babcock milk test (17.5 cc.) are measured in, and the test continued in the regular manner employed for milk. The reading of

MILK. 195

the fat is multiplied by 18 and the product, divided by the weight of cream taken, gives the per cent of fat.

U. S. Standards.*—Standard Cream is cream containing not less than 18% of milk fat.

Standard Evaporated Cream is cream from which a considerable portion of water has been evaporated.

Adulteration of Cream.—In some localities fat standards are fixed for cream both "heavy" and "light," those falling below such standards being deemed adulterated.

The same preservatives are employed in cream as in milk, and are

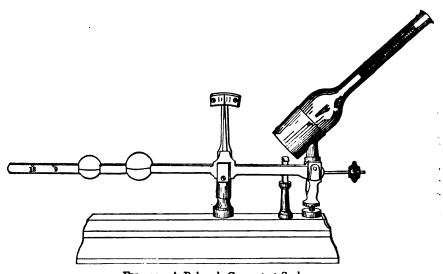


Fig. 55.—A Babcock Cream-test Scale.

detected in the same way. The color reaction for formaldehyde, by heating with hydrochloric acid and ferric chloride, is not as delicate in the case of cream as of milk, by reason of the large amount of fat. Before making the test the sample is preferably diluted with an equal volume of water, the heating is done in a casserole as with milk, but finally pour into a test-tube, and observe the color of the aqueous layer.

Gelatin in Cream.—Gelatin has been found by the writer in a number of samples of Massachusetts cream, its use being to increase the consistency. It was possible in one instance to obtain a sample of the adulterant used in the form of a powder, which proved on analysis to be chiefly gelatin

^{*} U. S. Dept. of Agric., Off. of Sec., Circ. 19.

with a small mixture of boric acid, these ingredients serving the two-fold purpose of thickening and preserving the cream.

Gelatin is best detected in cream or milk by the method of Stokes.* He uses for reagents (1) acid nitrate of mercury, prepared by dissolving metallic mercury in twice its weight of concentrated nitric acid (sp. gr. 1.42) and diluting with twenty-five times its bulk of water, and (2) a saturated aqueous solution of picric acid.

To about 10 cc. of the cream add the same amount of the acid nitrate of mercury solution and 20 cc. of cold water. The mixture is shaken vigorously and allowed to rest for five minutes, after which it is filtered. If much gelatin is present, the filtrate will not be clear, but opalescent. To the whole or a part of the filtrate a few drops of the picric acid solution are added, and if gelatin be present in any considerable amount, a yellow precipitate is formed. Avoid an excess of acid nitrate of mercury, as this would cause a precipitate with picric acid.

If gelatin is present in small amount only, a cloudiness is produced, best seen against a dark background. In the absence of gelatin, the solution will remain perfectly clear after adding the picric acid. The reaction is delicate to 1 part of gelatin in 10,000 parts of milk or cream.

Sucrate of Lime in Milk and Cream.—In the process of pasteurizing cream, a process which is becoming more and more prevalent, the consistency becomes reduced, so that while the value of the cream is actually enhanced on account of its freedom from bacteria and its increased capacity for keeping, its apparent richness is impaired when compared with untreated cream of the same composition. To restore this reduced consistency, Babcock and Russell † have shown that sucrate of lime may be used to advantage. They have applied the term "viscogen" to this compound, and have suggested that cream so treated, in order to be sold legally, should be known by some distinctive trade name as "viscocream" or "pasteurized visco-cream."

"Viscogen" is prepared by dissolving 2½ parts by weight of cane sugar in 5 parts water, and adding thereto, after straining, 1 part of quicklime slaked in 3 parts water. After shaking and settling, the supernatant liquid is syphoned off and bottled for use. It will thicken cream, milk, or condensed milk. The amount recommended to be added to cream should be two-thirds of the amount found by experiment necessary to neutralize its acidity.

^{*} Analyst, 22, p. 320.

[†] Wisconsin Exp. Station Bulletin, 54.

Sucrate of lime in milk or cream is indicated by the presence of sucrose, in connection with an abnormally high alkalinity of ash and excessive calcium oxide. These tests are carried out as follows:

Lythgoe's Modification of Baier and Neuman's Test for Detecting Sucrose.*—To 25 cc. of milk or cream, add 10 cc. of a 5% solution of uranium acetate, shake well, allow to stand for 5 minutes, and filter. To 10 cc. of the clear filtrate (in the case of cream use the total filtrate, which will be less than 10 cc.) add a mixture of 2 cc. saturated ammonium molybdate and 8 cc. dilute hydrochloric acid (1 part 25% acid and 7 parts water), and place in a water-bath at a temperature of 80° C. for 5 minutes. If the sample contains sugar, the solution will be of a prussian blue color. This should always be compared in a colorimeter with the standard prussian blue solution, prepared by adding a few drops of potassium ferrocyanide to a solution of 1 cc. of 1% ferric chloride in 20 cc. of water.

It has been claimed that pure milk will give this test. Occasional samples of pure milk will give a pale blue color, but this can be entirely removed by filtration and the filtrate will be green, while the color due to sugar will pass through the filter, giving the usual blue solution characteristic of adulterated samples. The color produced is due to a reduction of the molybdic acid, and is produced by levulose and dextrose, as well as by sucrose. Solutions of 1 gram of lactose, levulose, dextrose, and sucrose in 35 cc. of water were used in comparing the amount of color produced when heated with the molybdenum reagent for 5 minutes. Lactose produced no color, levulose gave a heavy blue, sucrose a weaker blue, and dextrose the weakest blue, the colors of the last three corresponding in intensity as 10:3:1.

Stannous chloride and ferrous sulphate give this blue color, but the reaction takes place in the cold, and, in small quantities, the color disappears on heating. In order for the color to persist after heating, the sample of cream must contain these substances to the extent of 1%, calculated as the metal. In this case the sample will be completely coagulated, and the taste will be very disagreeable. Hydrogen sulphide will also give the blue color, but it will disappear on heating. If the solution does not show the blue color before heating, it is free from hydrogen sulphide, ferrous sulphate, and stannous chloride.

As a confirmatory test for sugar, the resorcine test may be applied

^{*} Zeits. Unters. Nahr. Genussm., 16, 1908, p. 51.

to the serum prepared with uranium as described above. This test is given by sucrose and levulose, but not by dextrose or lactose.

Determination of Alkalinity of Ash and Calcium Oxide. — Weigh 25 grams of cream into a platinum dish, place in an oven at about 125–150° C. over night, and burn to an ash in a muffle at a low-red heat. Dissolve the ash in 20 cc. N/10 sulphuric acid, boil to expel the carbon dioxide, and titrate back with N/10 sodium hydroxide, using phenolphthalein as the indicator. Express results as cc. N/10 acid required to neutralize the ash of 100 grams of cream.

Make the final solution of the above determination acid with acetic acid, heat to boiling, add I gram of sodium acetate, and to the clear solution add an excess of ammonium oxalate, boil for a few minutes, filter, and wash with water. Dissolve the calcium oxalate in hot dilute sulphuric acid, and titrate hot with N/IO potassium permanganate. The number of cubic centimeters of N/IO permanganate, multiplied by 0.0112 (4×0.0028), gives the percentage of CaO in the sample.

The alkalinity of the ash of 100 grams of a sample of cream treated by the writer with 5 cc. of viscogen per quart was found to be 18.8, in terms of N/10 acid, added in excess and titrated back with N/10 alkali, using phenolphthalein. The alkalinity of the ash of 100 grams of pure cream (7 samples) varied between 6.4 and 13.6. The calcium oxide and alkalinity of ash diminish as the fat increases. Cream samples treated with calcium sucrate, having a fat content from 26 to 33%, show as a rule an alkalinity of ash of from 14 to 18, and a CaO content of from 0.15 to 0.175%. Creams of the same class untreated show in general alkalinity of ash not exceeding 12.5 and a CaO content not exceeding 0.135.

With higher fat contents both constants drop. For example, a cream of 45% fat containing calcium sucrate had an alkalinity of ash of 10.2 and a CaO content of 0.12%. Cream of about 45% fat untreated had an ash alkalinity of 6.5 and a CaO content of 0.103%.

ICE CREAM.

For many years a wide variety of iced foods have been made and sold under the general name of ice cream, many of which are largely composed of ingredients other than milk or cream. In the study and classification of foods of such a miscellaneous nature as ice cream, in its popularly accepted meaning, it is not always easy to satisfactorily define

MILK. 199

and fix standards. Whether, for example, the product should consist exclusively of frozen cream, sugar and flavoring, or whether eggs and other materials should be allowed under the unqualified name of ice cream, is a subject of some controversy.

Properly speaking, many mixtures sold under the name should be otherwise designated, as, for example, "frozen custard," to specify more aptly their nature and composition. The following standards show the attitude of the government in this regard:

U. S. Standards.*—Ice cream is a frozen product made from cream and sugar with or without a natural flavoring, and contains not less than 14% of milk fat.

Fruit ice cream is a frozen product made from cream, sugar, and sound, clean, mature fruits, and contains not less than 12% of milk fat.

Nut ice cream is a frozen product made from cream, sugar, and sound, non-rancid nuts, and contains not less than 12% of milk fat.

Fillers or Stiffeners.—In the manufacture of commercial "ice cream" substances are frequently added to cause the product to hold stiff and keep its consistency for many hours after freezing. The thickeners or fillers most commonly thus used are starch, gelatin, and gums such as gum tragacanth. Agar-agar and commercial casein are also said to be employed for this purpose.

Preparations are on the market sold for thickening ice cream. consisting, as a rule, of one or more of the above-named substances.

METHODS OF ANALYSIS.

Fat.—Howard's Method.†—With a wide-mouthed, free-flowing pipette, weigh 18 grams of the freshly melted and mixed sample into a Babcock cream bottle. Add 3 cc. of chloroform, and water to about three-fourths the capacity of the bottle. Shake, add 10 cc. of Fehling copper solution, shake again, and whirl for three minutes in the centrifuge. The fat is confined to the underlying chloroform layer. If the supernatant liquid is turbid, gum tragacanth or gelatin is probably present, in which case add 2 or 3 cc. of N/10 alkali, which causes these substances to precipitate.

Remove most of the supernatant liquid by aspiration through a narrow tube inserted in the cream bottle. Repeat the washing once, and then

^{*} U. S. Dept. of Agric., Office of Secretary, Cir. 19.

[†] Jour. Am. Chem. Soc,. 29, 1907, p. 1623.

blow in steam through a narrow tube till the chloroform is completely expelled. Cool, add water to a volume of 17.5 cc., and proceed as in the Babcock process for fat in milk.

Second Method.*—Nine grams of the sample are weighed into the test bottle, and 30 cc. of a mixture of equal parts by volume of concentrated hydrochloric acid and 80% acetic acid are added. Heat on the water-bath till the mixture darkens, but avoid charring. Whirl in the centrifuge, and read the percentage of fat directly, after adding hot water to run the fat layer into the neck. If charring has interfered with the fat reading, add ether after whirling to dissolve the fat layer, and draw off the ether solution into another bottle. Evaporate off the ether, fill with hot water, and again whirl and read.

Examination of the Fat.—Enough of the fat is usually obtained from its determination in the Babcock bottle, as above, to examine its character with the refractometer. Cottonseed and other oils are sometimes used as substitutes for milk fat. If a large amount of fat is needed, Howard suggests removing 30 to 40 cc. of the cream layer to a fat bottle, adding 1 cc. of acid nitrate of mercury and 20 cc. of petroleum ether, and whirling in a centrifuge, afterwards removing the ethereal layer, washing with water, and evaporating the ether.

Detection of Thickeners.—Patrick's Method.†—Add 25 cc. of water to 50 cc. of the sample, and boil till any thickener present is dissolved. Add 2 cc. of a 10% solution of acetic acid, heat to boiling, add 3 heaping teaspoonfuls of kieselguhr, and after shaking pass at once through a plaited filter. To 3 cc. of the clear filtrate add 12 cc. of 95% alcohol and mix thoroughly, thus precipitating the milk proteins not already removed, and also the gums and some of the gelatin, if much is present. Add 3 cc. of a mixture of 95 cc. of 95% alcohol and 5 cc. of concentrated hydrochloric acid. This acidified alcohol dissolves completely the milk proteins, and, if a clear solution then remains, no gums or vegetable jellies have been used as thickeners. Turbidity does not, however, necessarily indicate presence of a thickener, as it may be caused by a large amount of eggs, or by the souring of the ice cream. Dilute the mixture, if turbid, by adding 3 cc. of water. Any precipitate due to gelatin or eggs will be dissolved at this dilution, but not that due to vegetable gums. If gum tragacanth be present, the precipitate will be

^{*} Rep. Illinois State Food Commissioner, 1906, p. 82.

[†] U. S. Dept. of Agric., Bur. of Chem., Bul. 116, p. st.

MILK. 201

stringy and cohesive, especially after shaking, while agar-agar or other vegetable thickeners will cause a fine flocculent precipitate.

Souring of the ice cream sometimes produces a turbidity or precipitate under the above conditions, which is not always dissolved after diluting with water. Formation of such a precipitate (due to sourness) may, however, apparently be prevented by the previous addition of formaldehyde to the sample.

Howard's Test for Gums.—Precipitate 10 cc. of the melted sample with acetone, and wash with 2 or 3 portions of dilute alcohol, using the centrifuge. Boil the washed residue with 6 to 8 cc. of water and 1 cc. of 10% sodium hydroxide solution for half a minute. Cool, let stand a few minutes, filter, and heat the filtrate to boiling. Add one and one-half volumes of warm alcohol and shake. If agar-agar or gum tragacanth be present, a flocculent precipitate will immediately separate. Disregard a mere turbidity. To prove the absence of any considerable quantity of milk proteins in the precipitate, dissolve in cold water and saturate the solution with ammonium sulphate.

Gelatin.—Use the method of Stokes (p. 196) on 10 to 15 cc. of the sample, disregarding a faint cloudiness at the end.

Starch is detected by the usual iodine test.

Preservatives.—Formaldehyde and boric acid are tested for as in milk.

BUTTER.

The value of butter as a food depends almost entirely on its fat content, although minute quantities of protein and milk sugar are also included in its composition.

Hence butter is more logically treated in detail under the heading of fats (page 529).

CHEESE.

Nature and Composition.—Cheese consists principally of the curd and fat removed in a mass from milk, which has been curdled by the natural souring of the milk, or by the action of rennet. The separated mass of curd and fat, after being compressed, is allowed to undergo certain changes, which constitute the ripening or curing, due to the action of micro-organisms and enzymes. Sometimes cream is used as the source of cheese and sometimes skimmed milk. During the ripening process, which requires from a few weeks to several months, the characteristic

flavor is developed by the changes which the proteins undergo, and the digestibility of the cheese is improved. The nature of the proteolytic changes that take place during ripening are very little understood, but a variety, of complex nitrogenous products are formed, which Van Slyke divides as follows: paracasein, unsaturated paracasein lactate, paranuclein, caseoses (albumoses), peptones, amides, and ammonia. Besides nitrogenous bodies and fat, which are its chief constituents, cheese contains notable quantities of water, milk sugar, lactic acid, and mineral matter.

In some kinds of cheese salt and coloring matter are added.

Varieties.—Well-known cheeses of commerce are often named from districts, towns, or localities where they originated or are still made. They may be classified as cream, whole-milk, or skimmed-milk cheese, according to the quality of the product from which they are made, or again as hard, medium, or soft, according to whether (1) they are pressed, or (2) allowed to drain for days and sometimes weeks without pressure to a firm consistency, or (3) are made in the space of a few hours, being quickly drained on a sieve by hand pressure.

Cheddar Cheese, which is the common cheese of the United States (though originally made some 250 years ago in England and still made there), is a type of the hard cheese. Stilton, an English, and Gruyère, a Swiss cheese, belong to the medium class, and soft cheeses are represented by Brie and Neujchatel, both French cream cheeses. Other well-known varieties are Edam, a round, mild, long-keeping Dutch cheese, Camembert, a rich cream cheese, and Roquefort, made originally from ewe's milk in the French town of that name, and ripened in caves in the mountains. It is flavored by a peculiar mold.

The following table, compiled by Woll,* shows the average composition of various cheeses of commerce, both foreign and domestic:

	Water.	Casein.	Fat.	Sugar.	Ash.
	Per cent.	Per cent.	Per cent.	Per cent.	Per cent.
Cheddar	34.38	26.38	32.71	2.95	3.58
Cheshire	32.59	32.51	26.06	4-53	4.31
Stilton	30.35	28.85	35 - 39	1.59	3.83
Brie		17.18	25.12	1.94	5.41
Neufchatel	44-47	14.60	33.70	4.24	2.99
Roquefort	31.20	27.63	33.16	2.00	6.01
Edam	36.28	24.06	30.26	4.60	4.90
Swiss	35.80	24.44	37.40		2.36
Full cream, mean of 143 analyses	38.60	25-35	30.25	2.03	4.07

^{*} Dairy Calendar, p. 223.

Van Slyke has furnished the following analysis of the nitrogen compounds in a sample of normal American Cheddar cheese six months old and cured at 60° F.:

Per Cent	Per Cent	Per Cent N as Paracasein Mono- lactate.	Per Cent	Per Cent	Per Cent	Per Cent	Per Cent
N in	Water-		N as Para-	N as	N as	N as	N as
Cheese.	soluble N.		nuclein.	Caseoses.	Peptones.	Amides.	Ammonia.
3.80	1.46	0.94	0.14	0.22	0.18	0.79	0.13

U. S. Standards.*—Cheese is the sound, solid, and ripened product made from milk or cream by coagulating the casein thereof with rennet or lactic acid, with or without the addition of ripening ferments and seasoning, and contains, in the water-free substance, not less than 50% of milk fat. By act of Congress, approved June 6, 1896, cheese may also contain added coloring matter.

Skim-milk Cheese is defined the same as cheese except that it is made from skim milk, and no minimum percentage of fat in the water-free substance is specified.

Adulteration.—Cheese is commonly adulterated in two ways: first, by the partial or total substitution for the milk fat of a foreign fat, as oleomargarine or lard, and, second, by using skimmed milk as a material for its manufacture.

In many localities a standard percentage for fat in cheese is fixed by law, as in the case of the U. S. standard noted above, all samples falling below that standard, unless sold as skim-milk cheese, being deemed adulterated.

Some states have specific standards for varying grades of cheese, classified as to their fat content. Thus under the Pennsylvania law t cheese is divided into five grades, as follows:

Full-cream cheese must contain not less than 32% butter fat.

Three-fourths cream cheese must contain not less than 24% butter fat.

One-half cream cheese must contain not less than 16% butter fat. One-fourth cream cheese must contain not less than 8% butter fat. All cheese having less than 8% fat must be branded "Skimmed Cheese."

The term "filled cheese" is commonly applied to a product in which a foreign fat, as oleo oil or lard, has been used. Filled cheese is more

^{*} U. S. Dept. of Agric., Off. of Sec., Circ. 19.

[†] Penn. Laws, 1901, Act. 95, p. 128.

commonly found in localities where a carefully enforced fat-standard law prevails, but, in the absence of a standard for fat in cheese, the manufacturer can cheapen his product much more readily and conveniently by selling a skim-milk cheese in place of the whole-milk article, though not without producing a sensibly inferior product.

METHODS OF ANALYSIS.

Obtaining a Representative Sample.—Method of the A. O. A. C.*—By means of a cheese-trier remove, if possible, three cylindrical plugs from the cheese perpendicular to the surface and in length equal to about half the thickness of the cheese, one at the centre, one near the circumference, and one midway between the two. About one inch in length is cut off from each plug from the end having the rind, and this is discarded. The remaining portions of the plugs are then finely divided and mixed as intimately as possible.

In place of the plugs a narrow, wedge-shaped segment may be cut from the cheese, reaching from the circumference to the center, the portions near the rind being removed, and the remainder of the piece being finely divided and mixed as before. Analyses should immediately be begun after obtaining the sample.

Determination of Water.—Two or three grams of the sample are carefully weighed in a tared platinum dish, and dried to constant weight in an oven at 100° C. The loss of weight is reckoned as water.†

Determination of Ash.—Ignite the residue from the moisture determination at a low, red heat, cool in a desiccator, and weigh.

Determination of Fat.—Lythgoe's Modified Babcock Method.—Weigh accurately about 6 grams of the sample in a tared beaker. Add 10 cc. of boiling water, and stir with a rod till the cheese softens and an even emulsion is formed, preferably adding a few drops of strong ammonia to aid in the softening and emulsionizing, and keeping the beaker in hot water till the emulsion is tolerably complete and free from lumps.

If the sample is a full-cream cheese, which is usually evident from its taste and appearance, a Babcock cream-bottle is employed. The contents of the beaker, after cooling, are transferred to the test-bottle

^{*} U. S. Dept. of Agric., Bur. of Chem., Bul. 46, p. 55.

[†] Previously ignited sand or asbestos is recommended by some as an absorbent to be placed in the dish, but the writer gets better results in most cases directly as above.

as follows: Add to the beaker about half of the 17.6 cc. of sulphuric acid regularly used for the test, stir with the rod and pour carefully into the bottle, using the remainder of the acid in two portions for washing out the beaker. Finally proceed as in the regular Babcock test for milk. Multiply the fat reading by 18 and divide by the weight of the sample taken to obtain the per cent of fat.

Short's Method.*—Grind to a uniform powder 2 to 5 grams of the sample, and about twice its weight of anhydrous copper sulphate. Place a layer of anhydrous copper sulphate about 2 cm. thick on the bottom of the inner tube of a Johnson or Knorr extractor, add the ground mixture, and rinse the mortar first with a little anhydrous copper sulphate and finally with ether. Extract for 16 hours, evaporate the ether from the extraction-flask, and dry the fat in a boiling-water oven to constant weight.

Werner-Schmidt Method.—Boil 2 to 3 grams of the sample in the Werner-Schmidt tube (p. 139) with 5 cc. of water and 10 cc. of concentrated hydrochloric acid till, with constant shaking, all but the fat is dissolved. Cool, add 25 cc. of ether, and shake the tube well. Draw off as much as possible of the ether, after separation, in the usual manner, and extract with four or five additional portions of the solvent.

Distil off the ether from the combined extractions, and weigh the fat. Determination of Protein.—From 1 to 2 grams of the cheese are treated by the Gunning or Kjeldahl method, adding after partial digestion a piece of copper sulphate the size of a pea to aid in the conversion.† $N\times6.25=$ protein.

Separation and Determination of Nitrogen Compounds.—Methods of Van Slyke.—Twenty-five grams of the sample are mixed in a porcelain mortar with an equal volume of clear quartz sand. Transfer the mixture to a 450-cc. Erlenmeyer flask, add about 100 cc. of water at 50° C., and keep the temperature at 50° to 55° C. for half an hour with frequent shaking. Decant the liquid through an absorbent-cotton filter into a 500-cc. graduated flask. Treat the residue with repeated portions of 100 cc. each of water, heating, shaking, and decanting as above till the filtrate, or water extract, at room temperature amounts to just 500 cc. exclusive of the fat floating on top, and use aliquot parts of this water extract for the various determinations.

^{*} U. S. Dept. of Agric., Div. of Chem., Bul. 35, pp. 15, 17, 225.

[†] Van Slyke, N. Y. Exp. Station, Bulletin 215.

¹ N. Y. Exp. Station, Bulletin 215.

Water-soluble Nitrogen.—Determine the nitrogen by the Gunning method in 50 cc. of the above water extract, corresponding to 2.5 grams of cheese.

Nitrogen as Paranuclein.—Add 5 cc. of a 1% solution of hydrochloric acid to 100 cc. of the above water extract (corresponding to 5 grams of cheese), and keep the temperature at 50° to 55° till the separation is complete, as shown by a clear supernatant liquid. Filter, wash the precipitate with water, and determine the nitrogen therein by the Gunning method.

Nitrogen as Coagulable Protein.—Neutralize the filtrate from the preceding determination with dilute potassium hydroxide, and heat at the temperature of boiling water till the coagulum,* if any, settles completely. Filter, wash the precipitate, and determine the nitrogen therein.

Nitrogen as Caseoses.—Treat the filtrate from the preceding with 1 cc. of 50% sulphuric acid saturated with C. P. zinc sulphate, and warm to about 70° C. till the caseoses settle out completely. Cool, filter, wash with a saturated solution of zinc sulphate acidified with sulphuric acid, and determine the nitrogen in the precipitate.

Nitrogen as Amides and Peptones.—Place 100 cc. of the water extract of cheese in a 250-cc. graduated flask, add 1 gram of sodium chloride and a solution containing 12% of tannin, till the addition of a drop to the clear supernatant liquid does not further precipitate. Dilute to the 250-cc. mark, shake, pour upon a dry filter, and determine the nitrogen in 50 cc. of the filtrate, which gives the amount of nitrogen in the amido-acid and ammonia compounds. Deduct from this the amount of nitrogen as ammonia separately determined, and the difference is the amido-nitrogen.

Nitrogen as pertones is obtained by subtracting the sum of the amounts of nitrogen as paranuclein, coagulable proteins, caseoses, amido-bodies, and ammonia from the total nitrogen in the water extract.

Nitrogen as Ammonia.—Distil 100 cc. of the filtrate from the above tannin-salt precipitation into standardized acid, and titrate in the usual manner.

Nitrogen as Paracasein Lactate.—Treat the residue insoluble in water in obtaining the water extract, with several portions of a 5% solution of sodium chloride, to form a 500-cc. salt extract of the same, in an analogous manner to that employed in preparing the water extract. Determine the nitrogen in an aliquot part of this salt extract.

^{*} According to Van Slyke a precipitate at this point is rare in cheese.

MILK. 207

Determination of Lactic Acid.*—Add water to 10 grams of the cheese sample at 40° C. till the volume equals 105 cc. Shake and filter. Titrate 25 cc. of the filtrate (equivalent to 2.5 grams of cheese) with tenth-normal sodium hydroxide, using phenolphthalein as an indicator.

Each cubic centimeter of decinormal alkali is equivalent to 0.009 gram lactic acid.

Determination of Milk Sugar.—Boil 25 grams of finely divided cheese with two successive portions of about 100 cc. each of water, decant through a filter, and finally transfer the residue upon the filter and wash with hot water. Make up the entire aqueous extract thus obtained, when cold, to 250 cc., and determine the milk sugar by either Fehling method.

Detection of Foreign Fat.—The cheese fat, separated in the manner described below, is subjected to the various processes detailed under butter, in precisely the same way, the fat of cheese being identical with that of butter. The most ready means for judging its purity consists in determining the refraction with the butyro-refractometer, and the Reichert number.

Separation of the Fat for Examination.—Place a quantity, say 25 grams, of the finely divided sample in a large Erlenmeyer flask, add about 100 cc. of petroleum ether, cork the flask and allow it to stand for several hours with frequent shaking. Decant the petroleum ether through a filter, evaporate off the solvent by the aid of heat, and the residue will be found to consist of nearly pure fat.

Or, wrap a sufficient portion of the finely divided sample in a muslin cloth, place this in a dish, and heat on the water-bath. The fat which runs out is afterward filtered and dried at 100°.

Sufficient cheese fat may usually be obtained for the refractometer reading from the neck of the test-bottle, after completing the Babcock test, and, usually (except in the case of skimmed-milk cheese), for the Reichert number.

Detection of Skimmed-milk Cheese.—In a cream cheese the fat should greatly exceed the protein; in a whole-milk cheese the per cent of fat should at least equal that of the protein, and is generally in excess. If the fat is considerably less than the protein, the cheese has undoubtedly been made from skimmed milk. The following analyses, made in the writer's laboratory, illustrate these grades:

^{*} U. S. Dept. of Agric., Bureau of Chem., Bul. 46, p. 56.

Varieties of Cheese	Water.	Pat.	Protein.*	Ash.
Full cream (soft)	37.63	47.40	13.70	1.27
Whole milk (hard)	37.63 21.89	47.40 38.00	37.71	2.40
Whole milk (soft)	55.95	24.00	37.71 16.49	3.56
Skimmed milk (soft)	62.17	15.20	21.36	1.27
Centrifugally skimmed milk (soft)	72.80	2.00	23.52	1.68

^{*} By difference.

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CHAPTER VIII.

FLESH FOODS.

MEAT.

General Structure and Composition.—Meat is structurally made up of muscle fibers, held together by connective tissue, through which fat cells are usually more or less abundantly distributed. Each muscle fiber has a sheath or covering known as sarcolemma, formed of an albuminoid substance similar to elastin, and within the fibers are contained the meat juices, which are solutions in water of proteins, non-protein-nitrogenous extractives, and salts. The substance of the connective tissue is made up largely of the albuminoids elastin (insoluble) and collagen, the latter being convertible by boiling with water or treatment with acids into gelatin. The proteins of the meat juices consist chiefly of the globulin myosin (by far the most abundant), muscle albumin, and the muscle pigment hæmoglobin, or a substance closely analogous thereto.

In the living muscle there are no peptones, but the ferment pepsin is present. After death, by the action of the pepsin in presence of lactic acid, a portion of the normal proteins of the muscle undergoes, as it were, digestion, so that in meat both peptones and proteoses * are found.

The non-protein-nitrogenous extractives are mainly creatin, xanthin, hypoxanthin, and carnin, which, from their basic character, are known as flesh bases.

The approximate proportions in which the chief constituents are present in meat is thus shown by König:

Water	***************************************	75.0	to '	77.0
	Sarcolemma (muscle fiber)	13.0	to	18.0
	Connective tissue	2.0	to	5.0
	Albumin			
	Creatin	0.07	to	0.34
Nitrogenized compounds	HypoxanthinCreatinin	0.01	to	0.03
Millogenized compounds	Creatinin			_
	Xanthin	Undete	ermi	ined
	Inosinic acid			
	Uric acid			
	Urea	0.01	to	0.03
•				

^{*} A proteose or albumose known as myoalbumose normally exists in the live muscle.

Fat		0.5	to	3.5
	Lactic acid	0.05		
Other nitrogen-free compounds	Acetic acid	Undete	rmi	ned
Salts	Glycogen	(o.3	to	0.5) 1.8)
Composed o		(0.0	••	2.0,
-	Potash	0.40	to	0.50
	Soda		to	60.0
	Lime	0.01	to	0.07
	Magnesia	0.02	to	0.05
	Oxide of iron	0.003	to	0.01
	Phosphoric acid	0.40	to	0.50
	Sulphuric acid		to	0.04
	Chlorine	0.01	to	0.07

Nitrogen compounds constitute by far the most abundant and important portion of the substance of lean meat. Carbohydrates are almost entirely lacking, the small amount of glycogen and muscle sugar together constituting rarely more than 1 per cent.

Glycogen (C₆H₁₀O₅), sometimes called animal starch, is a white, amorphous, tasteless, and odorless substance, when pure, much resembling starch. It is soluble in water, forming an opalescent solution, and is insoluble in ether and alcohol. With iodine a port-wine color is produced, which disappears on heating and reappears on cooling. Glycogen is strongly dextro-rotary. It is converted to dextrose by boiling with dilute mineral acid.

Muscle Sugar is either entirely absent in the living muscle, or exists in traces only. After death it is formed presumably from the glycogen, and exists in a very minute quantity, probably as dextrose.

Inosite (C₀H₁₂O₀+ H₂O) is found in traces in the muscular substance of the heart, liver, kidneys, and testicles.

Proximate Constituents of the Commoner Meats.—The chief characteristics of the flesh of various animals are in the main very similar, whatever the nature of the animal. So true is this, indeed, that it is extremely difficult from a chemical analysis to distinguish a particular kind of flesh when mixed with that of other animals in finely divided meat preparations, such as sausages, potted and deviled meats, and the like.

The average composition of the commoner cuts of beef, veal, mutton, lamb, and pork, as well as of fowl and game, is shown in the following tables, compiled from the work of Atwater and Bryant,* the accompanying diagrams serving to locate, in the case of ordinary meats, the portion of the animal from which the meat is taken.

^{*}U. S. Dept. of Agric., Off. of Exp. Stations, Bul. 28 (Revised Ed.).

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6

FLESH FOODS.

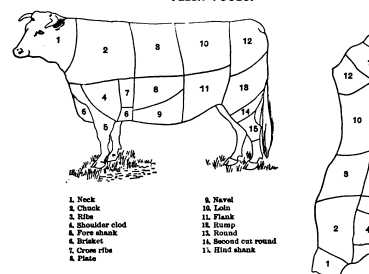


Fig. 56.—Diagram Showing Cuts of Beef. COMPOSITION OF BEEF.

			Num-		 	Pro	tein.			Fuel Value
	Cu	t.	yses.		Water.	N× 6.25.	By Differ- ence.	Fat.	Ash.	per Pound. Cals.
Chuck:	Lean-	edible portion	2		71.3	20.2	19.5	8.2	1.0	720
		as purchased	2	19.5	57-4	16.3	15.7	6.6	0.8	580
	Medium-	-edible portion	4		68.3	19.6	18.9	11.9	0.9	865
		as purchased	4	15.2	57.9	16.6	16.0	10.1	0.8	735
	Fat—	edible portion	4		62.3	18.5	18.0	18.8	0.9	1135
		as purchased	3	14-7	53-3	15.9	15.4	15.9	0.7	965
Ribs:	Lean—	edible portion	6		66.0	16.5	16.9	9.8	0.8	790
		as purchased	6	22.6	52.6	15.2	14.8	9-3	0.7	675
	Medium-	-edible portion	15		55-5	17.5	17.0	26.6	0.9	1450
	_	as purchased	15	20.8	43.8	13.9	13.5	21.2	0.7	1155
	Fat—	edible portion	9		48.5	15.0	15.2	35.6	0.7	1780
	_	as purchased	8	16.8	39.6	12.7	12.4	30.6	0.6	1525
Loin:	Lean—	edible portion	12	• • • • • •	67.0	19.7	19.3	12.7	1.0	900
		as purchased	11.	13.1	58.2	17.1	16.7	II.I	0.9	785
	Medium-	edible portion	32		60.6	18.5	18.2	20.2	1.0	1190
	_	as purchased	32	13.3	52.5	16.1	15.8	17.5	0.9	1040
	Fat—	edible portion	6		54-7	17.5	16.8	27.6	0.9	1490
_	_	as purchased	6	10.2	49.2	15.7	15.0	24.8	0.8	1305
Rump:	Lean—	edible portion	4		65.7	20.9	19.6	13.7	1.0	965
		as purchased	3	14.0	56.6	19.1	17-5	11.0	0.9	820
	Medium-	edible portion	10	[• • • • • ·	56.7	17.4	16.9	25.5	0.9	1400
	_	as purchased	10	20.7	45.0	13.8	13.4	20.2	0.7	1110
	Fat—	edible portion	5		47.1	16.8	16.4	35-7	0.8	1820
	-	as purchased	5	23.0	36.2	12.9	12.6	27.6	0.6	1405
Round:	Lean-	edible portion	31		70.0	21.3	21.0	7-9	1.1	730
	36 11	as purchased	29	8.1	64.4	19.5	19.2	7-3	1.0	670
	Medium-	edible portion	18		65.5	20.3	19.8	13.6	1.1	950
	TP-4	as purchased	14	7.2	60.7	19.0	18.3	12.8	1.0	895
	Fat-	edible portion	5		60.4	19.5	19.1	19.5	1.0	1185
		as purchased	3	12.0	54.0	17.5	17.1	16.1	0.8	1005
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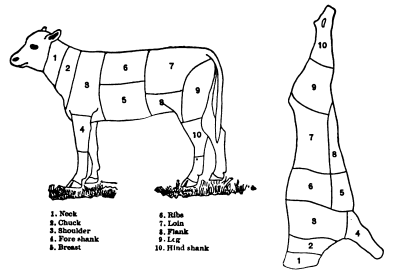


Fig. 57.—Diagram Showing Cuts of Veal.

COMPOSITION OF VEAL.

			Num-			Pro	tein.			Fuel
	Cu	t.	ber of Anal- yses.		Water.	N × 6.25.	By Differ- ence.	Fat.	Ash.	Value per Pound. Cals.
Chuck:	Lean-	edible portion	I		76.3		20.6	1.9	1.2	465
		as purchased	1	19.0	61.8		16.7	1.6	0.9	380
	Medium-	edible portion	6		73-3	19.7	19.2	6.5	1.0	640
		as purchased	6	18.9	59-5	16.0	15.6	5.2	0.8	515
Ribs:	Medium-	edible portion	9		72.7	20.7	20.1	6.1	1.1	640
		as purchased	9	25.3	54.3	15.5	15.0	4.6	0.8	480
	Fat-	edible portion	3		60.9	18.7	18.8	19.3	1.0	1160
		as purchased	3	24.3	46.2	14.2	14.2	14.5	0.8	875
Loin:	Lean—	edible portion	5		73-3	20.4	19.9	5.6	1.2	615
		as purchased	5 6	22.0	57.1	15.9	15.6	4.4	0.9	480
	Medium-	edible portion			69.0	19.9	19.2	10.8	10	825
	_	as purchased	6	16.5	57.6	16.6	16.0	9.0	0.9	690
	Fat—	edible portion	2		61.6	18.7	18.5	18.9	1.0	1145
_	_	as purchased	2	18.3	50.4	15.3	15.1	15.4	0.8	935
Leg:	Lean	edible portion	9		73-5	21.3	21.2	4.I	1.2	570
		as purchased	9	9.1	66.8	19.4	19.3	3-7	1.1	520
	Medium-	edible portion	10	 	70.0	20.2	19.8	9.0	1.2	755
		as purchased	9	14.2	60.1	15-5	16.9	7.9	0.9	620

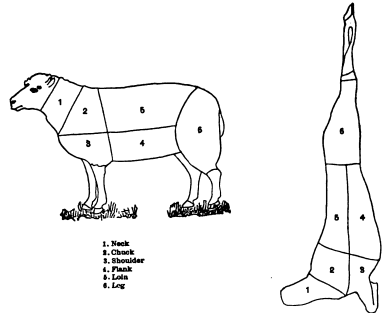
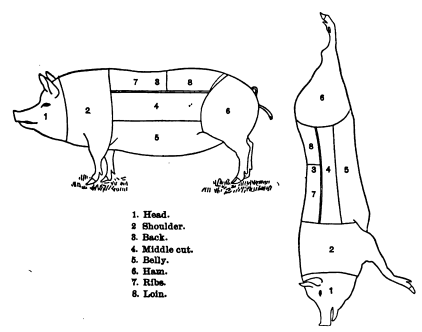


Fig. 58.—Diagram Showing Cuts of Mutton. COMPOSITION OF MUTTON AND LAMB.

			Num-			Pro	tein.			Fuel Value
	Cu	it.	ber of Anal- yses.	Refuse.	Water.	N× 6.25.	By Differ- ence.	Fat.	Ash.	per Pound. Cals.
	Мит	TON.								
Chuck:	Lean-	edible portion	1		64.7	17.8	18.1	16.3	0.0	1020
		as purchased	1	19.5	52.1	14.3	14.5	13.1	0.8	820
	Medium-	-edible portion	6		50.9	15.1	14.6	33.6	0.9	1700
		as purchased	6	21.3	39.9	11.9	11.5	26.7	0.6	1350
	Fat—	edible portion	2		40.6	13.9	13.7	44.9	0.8	2155
		as purchased	2	16.5	33.8	11.6	11.5	37-5	0.7	1800
Loin:	Medium-	-edible portion	13		50.2	16.0	15.9	33.1	0.8	1695
		as purchased	12	16.0	42.0	13.5	13.0	28.3	0.7	1445
	Fat-	edible portion	3		43-3	14.7	14.2	41.7	0.8	2035
		as purchased	3 8	11.7	38.3	13.0	12.5	36.8	0.7	1795
Flank:	Medium-	edible portion	8		46.2	15.2	14.8	38.3	0.7	1900
		as purchased	2	9.9	39.0	13.8	13.6	36.9	0.6	1815
Leg:	Lean—	edible portion	3		67.4	19.8	19.1	12.4	1.1	890
		as purchased	3	16.8	56.1	16.5	15.9	10.3	0.9	740
	Medium-	edible portion	11		62.8	18.5	18.2	18.0	1.0	1105
	_	as purchased	11	18.4	51.2	15.1	14.9	14.7	0.8	900
	Lab			i	_					
Chuck:		edible portion	I		56.2	19.1	19.2	23.6	1.0	1350
_		as purchased	I	19.1	45.5	15.4	15.5	19.1	0.8	1090
Leg:	Medium-	edible portion	2	¦	63.9	19.2	18.5	16.5	1.1	1055
	- .	as purchased	2	17-4	52.9	15.9	15.2	13.6	0.9	870
	Fat-	edible portion	1	•••••	54.6	18.3	17.1	27.4	0.9	1495
.		as purchased	I	13-4	47 - 3	15.8	14.8	23.7	0.8	1295
Loin:		edible portion	4	ا - یا	53-1	18.7	17.6	28.3	1.0	1540
		as purchased	4	14.8	45-3	16.0	15.0	24.I	0.8	1315



Fro. 59.—Diagram Showing Cuts of Pork.

COMPOSITION OF PORK, POULTRY, AND GAME.

			Num-			Pro	tein.			Fuel Value
	C	ut.	ber of Anal- yses.	Refuse.	Water.	N× 6.25.	By Differ- ence.	Fat.	Ash.	per Pound Cals.
	Po	RK.								
Should	er:	edible portion	19		51.2	13.3	13.8	34.2	0.8	1690
		as purchased	19	12.4	44-9	12.0	12.2	29.8	0.7	1480
Loin:	Lean-	edible portion	I		60.3	20.3	19.7	19.0	1.0	1180
		as purchased	1	23.5	46.I	15-5	15.1	14.5	0.8	900
	Fat—	edible portion	4	• • • • • • •	41.8	14.5	13.1	44.4	0.7	2145
	_	as purchased	4	16.5	34.8	11.9	10.9	37-2	0.6	1790
Ham:	Lean-	edible portion	2		60.0	25.0	24.3	14.4	1.3	1075
	_	as purchased	2	0.9	59-4	24.8	24.2	14.2	1.3	1000
	Fat-	edible portion	5		38.7	12.4	10.0	50.0	0.1	2345
		as purchased	5	13.2	33.6	10.7	9.2	43.5	0.5	2035
]	POULTRY A	ND GAME.		i '			1			
Chicker	n:	edible portion	3		74.8	21.5	21.6	2.5	1.1	505
		as purchased	3	41.6	43.7	12.Š	12.6	1.4	0.7	295
Fowl:		edible portion	26		63.7	19.3	19.0	16.3	1.0	1045
		as purchased	26	25.9	47-I	13.7	14.0	12.3	0.7	775
Goose:		edible portion	I		46.7	16.3	16.3	36.2	0.8	1830
		as purchased	I	17.6	38.5	13.4	13.4	29.8	0.7	1505
Turkey	:	edible portion	3		55-5	21.1	20.6	22.9	1.0	1360
•		as purchased	3	22.7	42.4	16.1	15.7	18.4	0.8	1075
Quail:		as purchased	ī		66.9	21.8		8.0	1.7	775
-			l	1			1 1			

Characteristics of Sound Meat.—The reaction of meat should be acid. If neutral or alkaline, decomposition is indicated, except that alkalinity may be due to the use of alkaline salts as preservatives.

Letheby* gives the following characteristics of sound, fresh meat. In color it is neither pale pink nor deep purple, the former indicating that the animal was affected with some disease, and the latter that it died a natural death, and was not slaughtered. In appearance it is marbled, due to the presence of small veins of fat distributed among the muscles. In consistency it is firm and elastic to the touch, and should hardly moisten the finger; a wet, sodden, or flabby consistency with a jelly-like fat is indicative of bad meat. As to odor, it is practically free; whatever odor there is should not be disagreeable; a sickly or cadaverous smell is indicative of diseased meat. After standing for a day or so, it should not become wet, but on the contrary should grow drier. When dried at 100° C. it should not lose more than 70 to 74 per cent in weight; unsound meat frequently loses 80% or more. It should shrink very little in cooling.

Inspection of Meat.—While carefully drawn laws exist almost everywhere relating to the sale of meat, and government inspectors are appointed to carry out the requirements of the laws, yet in this country there is undoubtedly some meat unfit for food on the market, owing to the small number of inspectors, and the consequent comparative safety with which unscrupulous dealers may sell meats forbidden by law and escape detection. The inspection of meats and fish under municipal ordinances is not always carried out as thoroughly as might be desired.

Unwholesomeness of Meat may be due to a diseased condition of the animal while alive, or to poisonous or injurious toxins developed by the action of bacteria after death. In the first case, the diseased conditions may be due to temporary causes only, or to the presence of animal parasites, such as trichinæ in pork, or as the result of pathogenic bacteria, causing such serious diseases as tuberculosis, anthrax, glanders, etc. It thus requires much skill and judgment on the part of the meat-inspector to correctly pass upon the suitability for food of the various meats as they appear on the market. Coplin and Bevan † give in detail useful data regarding the inspection of meat, as well as of the animal before slaughtering, showing the requisite size, weight, age, conditions of health etc., that should obtain. The detailed physical and microscopical examination involved in such inspection is, however, rarely germane to the work of the public food analyst, and will not be treated of in this manual.

^{*}Lectures on Food, p. 210.

It is also beyond the scope of the present work to treat of the harmful toxins developed by bacterial action in meat and fish, causing what is known as ptomaine poisoning. The work of detecting and isolating such poisons comes within the province of the bacteriologist and biologist, rather than that of the chemist, involving many experiments upon guinea-pigs, rabbits, or other animals not usually found in the chemist's laboratory. It has furthermore been recently shown by Vaughn and Novy * that even when these toxins are present in foods in sufficient quantity to produce serious results, very considerable amounts of the food must be taken in order to isolate them by chemical means, more, in fact, than is usually available for analysis.

For the general inspection of meats for animal parasites, poisonous toxins, etc., the reader is referred to such works as those of Vaughn and Novy, Fischöder, Walley, Andrews, Cobbold, and Salmon as cited in the references on pages 258 to 260.

U. S. Standards.†—Standard Meat is any sound, dressed, and properly prepared edible part of animals in good health at the time of slaughter. The term "animals" as herein used includes not only mammals, but fish, fowl, crustaceans, mollusks, and all other animals used as food.

Standard Fresh Meat is meat from animals recently slaughtered, or preserved only by refrigeration.

Standard Salted, Pickled, and Smoked Meats are unmixed meats preserved by salt, sugar, vinegar, spices, or smoke, singly or in combination, whether in bulk or in packages.

Standard Manujactured Meats are meats not included in the above divisions, whether simple or mixed, whole or comminuted, with or without the addition of salt, sugar, vinegar, spices, smoke, oils, or rendered fat, if they bear names descriptive of their composition, and when bearing such descriptive names, if force or flavoring meats are used, the kind and quantity thereof are made known.

Preservation of Meat.—Raw meat soon begins to decompose, unless precautions are taken to destroy, or at least check the growth of putrefying bacteria. From earliest times the subjection of meat to extreme cold has been practiced in order to enhance its keeping qualities. Bacterial growth is inhibited to a greater or less extent by refrigeration, by subjecting the meat to the various processes of curing, by the use of high temperatures and the exclusion of air as in canning, and by the employment of antiseptics.

^{*} Cellular Toxines.

[†] U. S. Dept. of Agric., Off. of Sec., Circ. No. 19.

Refrigeration may consist (1) in actually freezing the meat, in which condition it may be kept without decomposition almost indefinitely, until finally thawed for use, or (2) in keeping the meat at or near the temperature of freezing without actually congealing it, as is done by the use of the ordinary refrigerator. The second method, while much less efficacious than the first, serves to prevent decomposition for a considerable time and is preferred for beef, mutton, and pork. The lower temperatures are employed with poultry and game.

Curing consists in subjecting the meat to various processes of drying, smoking, pickling, corning, etc., or to a combination of these processes. In simple drying, the meat is subjected to the heat of the sun or to artificial heat. In smoking, which is commonly practiced on beef and ham, the meat, which may or may not be first salted or otherwise treated, is exposed for some time to the smoke of burning beech or hickory wood, during which it becomes to some extent impregnated with the antiseptic properties of the creosote and pyroligneous acid, at the same time being dried by the heat of the burning wood. In some cases best results are obtained by a slow smoking at a comparatively low temperature, while in others quick, hot smoking is found most efficacious. The character of the meat is decidedly changed by smoking, and, according to Utescher, smoked meat is always alkaline in reaction. In pickling, the meat may be treated with dry salt and subjected to pressure, so that the meat juice forms the liquid for the brine, in which it is allowed to remain for some time; or, as in the ordinary process of corning, the beef is soaked for some days in a strong solution of salt to which a little saltpetre (KNO₃) has been added. In the process of pickling, the salts from the brine slowly diffuse into the interior of the meat by osmosis, a part of the soluble albumin passing out into the brine. The effect of the saltpetre is to preserve the natural red color of the meat, which by the action of salt alone becomes destroyed, or at least impaired.

Bacon and ham are frequently cured by pickling in brine containing salt, saltpetre, and cane sugar, and sometimes also such antiseptics as boric acid and calcium bisulphite.

The curing of bacon is sometimes effected by injecting the pickling fluid into the tissues with a "pickle-pump," capable of exerting a pressure of 40 lbs. to the square inch, and provided with a hollow, perforated needle-nozzle, which penetrates the flesh. After pickling, the bacon or ham

may be simply dried, or, if desired, smoked. Oak sawdust is frequently burned for the smoking of ham.

The Use of Antiseptics in Meat.—Most of what might be termed the modern preservatives are to be looked for in one or another of the various meat preparations, though some are better adapted than others for use in particular cases, as will be seen by reference to the composition of typical commercial preservative mixtures given on page 817.

Borax and boric acid, usually in mixture, have been used more commonly than any other preservatives for the preservation of meat. Like salt, they are used commonly in surface application, in the case of large cuts of meat, or by mixing, in the case of sausage meat. A more recent method of application consists in impregnating the tissue of the meat with a solution of the boric mixture, by means of the above-described pickle-pump. The use of boric acid and its compounds, however, is not permitted under the regulations of the Federal meat inspection law of the United States and Germany.

Sulphurous Acid.—As much as 1% of a solution of sulphurous acid may be added to meat without being apparent to the taste or smell. Mitchell quotes Fischer as having found that 50% of the preserved meat products (sausages, etc.) sold in Breslau in 1895 contained sulphites, varying in amount from 0.01 to 0.34 per cent of sulphur dioxide. Calcium bisulphite is a salt commonly employed. In Hamburg steak it serves partly as a preservative, but chiefly as a deodorizer and a restorer of the bright red color of fresh meat.

Salicylic Acid is not of such common occurrence in meat products as the other antiseptics mentioned. The writer has found it in prepared mince-meat.

Among other preservative substances sometimes used with meat are solutions containing phosphoric acid and aluminum salts.

The toxic effects of these and other antiseptic chemicals in meats, and the most practical means of controlling their use are questions in controversy, presenting no new phases that have not been elsewhere discussed in treating of the general question of preservatives in food. Methods of detecting preservatives in meats are given elsewhere.

Effect of Cooking on Meat.—The general result of cooking is to render the meat less tough, to develop an agreeable flavor, and to coagulate more or less of the proteins. When subjected to moist heat, such as boiling and steaming, some of the soluble materials are dissolved, so that when the liquor in which the meat is boiled is thrown away, some of the valuable substances are lost. This is especially true when meat is placed in cold water which is afterwards brought to boiling, a method to be recommended when the liquor with the dissolved extractives is to be used for broth. When the meat to be boiled is placed at once in boiling water, there is less loss of soluble matter by reason of the formation of a more or less impenetrable coating on the outside, by the coagulation of the proteins. Meat that is boiled becomes softer, owing to a partial dissolving of the gelatin formed. In the dry cooking of meat, as by broiling or roasting, there is usually a hardening of the tissues, and a driving out of some of the meat juices, which are, however, often recovered in the form of gravies.

Canning of Meat.—By far the most effective method of preserving meat and meat preparations of all kinds for long periods of time, consists in the application of the principle of sterilizing by heat, and sealing in air-tight cans. The process of canning cooked meat and its products does not differ materially from that employed in the similar preparation of vegetables. (See Chapter XXI.) Previous to canning, the meats are usually cooked by boiling, during which process the changes described in the preceding paragraph take place.

The practice of misbranding chopped meat with respect to variety has been very prevalent in the past, and many varieties of so-called potted and devilled meats and game have frequently consisted wholly or in large part of a cheaper variety than that specified on the label. This practice has been largely corrected in this country, owing to the enforcement of the regulations of the Federal meat inspection law.

It is largely among the canned meats and prepared meat products that instances of adulteration are to be found, since the fresh meats in whole cuts are rarely subject to adulteration.

Preservatives are sometimes added to canned meats, especially in the case of dried and smoked beef, ham and bacon, and in the potted and devilled mixtures. Boric acid, benzoic acid, and sulphites have been found in these preparations.

It is believed, however, that this practice has been largely discontinued, owing to the enforcement of the Federal regulations mentioned above.

Composition of Canned Meats.—The following table, compiled from results published by Bigelow and others,* shows the composition of various of the most common canned and preserved meats and meat

^{*} U. S. Dept. of Agric., Bur. of Chem., Bulletin 13, part 10.

COMPOSITION OF CANNED MEATS.

) June	01110	COMI USITION OF CANNED MEATS.	בעוניי	יייי ריי	. CT U							
						Nitrogen.	gen.		Nitr	snouelo	Nitrogenous Substances.	.890		
		Number of Analy.	Water.	Fat.	Total.	Insolu- ble in Hot Water.	Precipi- tated by Bro- mine.	Meat Bases.	Protein N× 6.25.	Pro- teins Insolu- ble in Hot Water (Coag.).	Gela- tinoids & Pro- teins Ppt'd by Bro- mine.	Meat Bacce.	Total Ash.	Sodium Chlo- ride.
Fresh beef:	Average	2	68.19	12.60	3.02	2.41	61.0	0.43	18.89	15.08	1.21	1.29	96.0	0.08
	Maximum	:	71.17	15.33	3.19	2.52	0.24	0.53	19-94	15.75	0.50	1.65	1.13	0.24
Canned beef, roast and boiled:	Average	22	. &.	13.99	4.15	20.5	0.57	5. 54	25.95	19.29	3.59	1.58		0.53
	Maximum		66.39	31.78	5.51	4.4	1.16	1.03	4.6	27.94	7.25	3.21	3.51	2.51
Canned beef, corned:	Average	11	56.33	11.43	6.4	3.58	0.33	0.36	26.63	22.37	2.03	11.11	8%	3.37
	Minimum.		46.94	6.33	3.72	3.01	200	0.30	23.25	18.81	1.25	0.62	3.57	2.56
Canned beef, smoked and dried:	Average	11	47.42	7-46	5.21	4.22	0.15	0.85	32.59	26.41	0.92	2.63	12.51	6.67
	Minimum.		30.22	4.50	9. 4. 9. 8.	3.30	. %		25.50	20.05	.385	1.59	5.50	7.15
Fresh horse meat:	Average	91	69.8I	9.61	3.11	2.37	0.30	0.55	19.47	14.83	1.23	1.70	1.01	0.0
	Maximum		76.91	33.66	3.60	2.97	0.30	I.22	22.50	18.56	2.25	3.81	1.27	60°E
Canned ham and bacon:	Average	13	36.77	37.81	3.11	2.30	0.30	0.54	19.43	14.86	1.22	1.67	.88	5.34
	Maximum	•	53.30	69.07	88.	4.75	0.74	1.38	41.75	29.69	4.62	4.31	8.6	17.84
Canned tongue:	Average	17	15.34	20.23	3.11	2.50	0.21	0.00	10.43	15.64	1.33	1.23	3.71	2.00
)	Maximum	. :	71.80	38.00	3.95	3.20	0.48	0.72	24.69	20.00	3.8	2.25	6.22	4.43
Canned fowl:	Minimum	:	39.58	6.84	2.00	1.70	0.03	70.0	13.00	8 5	0.12	0.22	87.5	0.37
	Maximum	?	90.00	30.31	5.14	4.45	0.76	0.46	32.12	27.81	4.75	1.44	2.61	9.5
•	Minimum	:	47.46		2.14	1.77	0.11			11.06	: :		0.58	Trace
Canned towl and game:	Average	15	62.44		3.62	2.86	0.26			18.39	99.1	1.39	2.31	1.16
	Maximum	:	8:0	30.97	5-10	4.35	0.30	0.03	31.87	27.19	2.44	2.03	3.82	2.23
Canned sausages:	Average		2.5 2.5		2.22	1.20	0.12	0.10	12.00	7.50	0-75	0.00	8.6	0.37
	Maximum	:		41.34	3.86	3.86	0.62	0.42		20.75	3.88	1.31	6.37	. 6
	Minimum		44.02	3-53	0.52	0.38	0.03	0.04	3.25	2.38	0. ro	0.12	1.35	0.04

products, and in one or two instances fresh meat has been included for comparison.

Sausages.—Nature and Composition.—Sausages are made from finely chopped meat, highly seasoned with various spices, and, as usually sold, stuffed into casings made of the cleaned and prepared intestine-skin of cattle, sheep, or hogs. The meat most commonly used is pork. Sausages are frequently home-made, especially in farm communities, the chopped and seasoned meat being stuffed in cloth bags instead of casings. Any and all kinds of meat are used in sausages, and much that is undesirable and even unwholesome, is undoubtedly most readily used up in this product. There is little doubt that horse meat occasionally gets into the hands of the marketmen to be worked up in the form of sausages mixed with other meat. The condition in respect to these matters has been greatly improved, however, by the increased vigilance of State and Federal authorities. Sausages are sometimes artificially colored, and in some cases contain so-called "fillers" in the nature of dried bread, corn meal, potato starch, crackers, waste biscuit, boiled rice, etc.

CHEMICAL	COMPOSITION	OF	CATICACTC #

				Pro	tein.		Total		
Kind.	No. of Analy- ses.	Ref- use.	Water.	N×6.25	By Differ- ence.	Pat.	Carbo- hy- drates.	Ash.	Fuel Value, Cals.
Farmer: edible portion as purchased Pork: as purchased Bologna: edible portion	1 1 11 8	3.9	23.2 22.2 39.8 60.0	29.0 27.9 13.0 18.7	27.2 26.2 12.7 18.4	42.0 40.4 44.2 17.6	1.1 0.3	7.6 7.3 2.2 3.7	2310 2225 2125 1095
as purchased Frankfort: as purchased	4 8	3-3	55-2 57-2	18.2	18.0	19.7	1.1	3.8 3.4	1170

^{*} U. S. Dept of Agric., Off. of Exp. Stations, Bul. 28 (Revised Ed.).

Adulteration of Sausages with Starchy Materials and Water.—Robison, who has made a special study of these forms of adulteration at the Michigan Dairy and Food Department, states as follows:* "Lean meat carefully chopped has an enormous combining power and can be made to take up a great quantity of water. Frankfurts, bologna, and pork sausage have been found to be adulterated with from 0.5 to 5% of starch, indicating an addition of approximately 1 to 10% of so-called cereal (chiefly corn flour), and from 5 to 40% of water in addition to

^{*} Personal communication.

that contained in the meats when in their fresh condition. The main excuse for the use of water is that it renders the meat of such a consistency that it may be easily stuffed into thin cases, such as are usually used for sausages that are eaten without removing the casing. As a matter of fact, this addition is not necessary where fresh meats are used, nor with those cuts of meat which the American public is in the habit of using in the manufacture of sausages in the home. Without doubt, in sausages composed of ox hearts, ears, snouts, lips, etc., in considerable quantities, the addition of water may facilitate the stuffing into thin casings.

"Starch hastens and increases the absorbing or combining power of lean meat. In many instances where inferior products, such as ears, etc., are used, virtually it is the only absorbing agent present in the product. It then serves a two-fold purpose, first, giving an absorbing power to meat which it has not, or inflating the absorbing power of a meat which naturally is deficient in this respect, and second, acting as a skeleton or framework, thereby disguising shrinkage during the process of cooking. Generally, added water and cereal are evidences of inferiority, and they are by no means infrequently added with the very purpose of concealing such inferiority.

"The evidence of adulteration with water is the discrepancy in the ratio of the water to the protein in the sausage. This ratio in sausage made from the fresh carcass varies from 3:1 to 3.6:1, being on an average about 3.35:1."

Artificial Coloring Matter in Sausages.—Owing to the rapid color changes which freshly chopped meat, especially beef and mutton, naturally undergo, it is a common practice to employ powdered niter or salt-peter. Treated in this manner, meat remains pink, owing to the action on the hæmoglobin of the oxides of nitrogen resulting from the nitrate. As much as 4 ounces of niter to 100 lbs. of meat is sometimes used. A larger quantity would result in a shriveled appearance. The use of artificial colors has been common in the past, in order to permanently dye the flesh a bright red, similar to the tint which the oxy-hæmoglobin naturally imparts to the beef when fresh. A variety of colors have been employed for this purpose, such as red ocher, coal-tar dyes, cochineal, etc. They were sometimes used in admixture with preservatives. Their use has been largely discontinued in this country, owing to the enforcement of the regulations under the Federal meat inspection law.

ANALYTICAL METHODS.

In analyzing meats and meat products due regard must be paid to their perishable nature, and, for this reason, immediately after their receipt by the analyst the various determinations should be promptly begun and rapidly carried out. If delays are absolutely necessary, the samples, as well as some of the solutions, especially during the earlier course of the analysis, should be kept on ice to prevent decomposition. Even at low temperatures, however, both bacterial and enzymic decomposition occur, and the nature of the proteins is slowly changed. Refuse material, such as bones, skin, gristle, tendons, etc., are separated as completely as possible by means of a knife from the edible portion, and the latter, cut first into small pieces, is passed repeatedly through a sausagemachine or ordinary household meat-chopper, in order to reduce to a homogeneous, finely divided mass.

Determination of Water. —From 1 to 3 grams of the finely divided material are weighed in a tared platinum dish, and dried to minimum weight at a temperature of 100° C. in an air-oven. A slight oxidation of the fat may introduce a trifling error, but, excepting for the most exact work, where the drying should be accomplished in an atmosphere of hydrogen, or *in vacuo*, the above method is sufficiently close.

Determination of Water in Sausages.—Robison's Method.—A large sample (100 to 500 grams) is put through a food-chopper, weighed on a large porcelain plate, and allowed to dry at 70 to 90° C. A steam radiator may be conveniently used for this purpose. After drying 10 to 12 hours, or over night, it is reweighed and finely ground in a small laboratory mill. If the sample is quite fat, the preliminary drying of the chopped meat may be carried out conveniently on a sieve, which will permit the fat to drain through onto a plate below, thereby making more simple and accurate the sampling and mixing. The fat thus removed should be separately weighed and dried. If the sample is quite lean, the final drying of 2 to 5 grams of the air-dried sample may be made at 100° C. in an ordinary water or electric oven. If it is quite fat, it is best to conduct this drying in a current of hydrogen.

Determination of Ash.—The residue from the total solids is incinerated in the original dish over the free flame at a low red heat, or in a muffle. It is usually advantageous, especially in the case of salt meat, to exhaust the charred sample with water, and collect the insoluble residue on a filter and ignite. The filtrate is then added, evaporated to dryness, and

the whole heated to low redness and weighed. A perfectly white ash is difficult to obtain.

Determination of Fat.—Extraction Method.—About two grams of the sample are carefully weighed in a tared Schleicher-and-Schüll Soxhlet shell, which, after drying in the air-oven at 100°, is transferred to an extraction apparatus and subjected to continuous extraction for at least sixteen hours with anhydrous ether, or pure petroleum ether. It is impossible to extract all the fat in this manner, but the approximate result obtained is as a rule accepted, since complete extraction involves digestion of the nitrogenous matter with pepsin, and intermittent treatment with the solvent, a process both tedious and open to other sources of error.

More complete results may be obtained by pulverizing the extracted residue in a glazed porcelain dish with a glazed pestle, transferring again to the extraction shell, rinsing the dish with ether or petroleum ether, and again extracting.

Kita's Centrifugal Method.*—Two and one-half grams of meat are treated in a Babcock milk flask with 8 cc. 1:1H₂SO₄ (or 5 grams with 17 cc.), and heated in a water bath to 60 to 70° with occasional agitation till the solution of the proteins is complete. One cc. of amyl alcohol is then added, and sufficient dilute H₂SO₄ to bring the layer of fat within the graduated scale. The tube is then whirled in a centrifuge for from 3 to 5 minutes, and the amount of fat read on the scale. The amyl alcohol may sometimes be dispensed with, but is usually necessary for complete extraction and to obtain a clear layer of fat.

Examination of Fat.—In case a special examination of the fat is to be made, a large portion of the original finely divided sample is shaken in a corked flask with petroleum ether boiling below 60°, and allowed to digest for some hours or over night. The solvent is then poured off, the petroleum ether removed as far as practicable by distillation, and last traces of the solvent removed by allowing the fat to stand in a vacuum desiccator over freshly ignited calcium chloride. In the case of mixed canned-meat preparations, it is often desirable to determine the character of the fats as a possible clue to the variety of meat used. For this purpose the regular methods prescribed under oils and fats (Chapter XIII) are used.

Determination of Total Nitrogen.—Two grams of the sample are treated according to the Gunning or Kjeldahl method (page 69). While in the case of meat the time-honored custom of representing the total protein or nitrogenous substances by $N \times 6.25$ is by no means strictly

^{*} Arch. f. Hyg., 51, pp. 165-78.

FATS OF VARIOUS MEATS AND MEAT PREPARATIONS.*

	FAIS OF VARIOUS MEALS AND MEAL FREFARATIONS.	ARIOD	MEA	מאת כו	MEA	LALL	7110	ING.				
		Specific Gravity 100° 100°	Specific Gravity 100° 15°.	Degrees Butyro- refrac- tometer at 35°.	Index of Refrac- Ition at 35°.	Melting-	Melting-Chilling- point. point.	Iodine Number	Koetts- torfer Number.	Soluble Acids.	In- soluble Acids.	Heat with H ₂ SO ₄ .
Beef, roast and boiled:	Average Maximum	0.8953 0.9046 0.8925	0.8589 0.8679 0.8563	52.8 55.5 47.0	1.4610 1.4629 1.4571	40.1 43.9 36.5	32.2 37.0 27.8	45.6 50.6 36.1	194.8 200.0 188.0	0.45	93.54 95.27 91.90	35.8 36.0 35.6
Beef, canned, corned:	Average Maximum Minimum	0.8934 0.8944 0.8925	0.8572 0.8581 0.8563	55.0	1.4618 1.4631 1.4609	40.1 43.4 37.2	31.0 34.5 29.9	42.7 48.6 37.9	196.3 210.0 191.0	0.60	92.37 94.50 89.51	36.2 37.0 35.5
Beef, canned, dried, and smoked:	Average Maximum			55.2 58.5 51.0	1.4627 1.4649 1.4599	39.3 41.6 37.7	26.0 29.0	54.1 57.5 50.9		-		
Horse meat:	Average Maximum Minimum	0.9067 0.9184 0.8868	0.8811 0.8508	62.3 76.5 55.2	I.4673 I.4762 I.4625	29.7 32.5 27.2	17.2 25.0 12.0	66.4 77.0 61.4	201 205 . 196	1.32 2.76 0.19	91.37 95:44 88.18	56.5 46.2
Ham and bacon, canned:	Average Maximum Minimum	0.8893	0.8533	53.4 58.2 49.0	1.4615 1.4648 1.4586	30.5	20.3 24.0 17.5	57.8 68.2 48.5	193 207 179	0.98 2.63 0.19	90.80 94.51 86.40	41.4 43.5 39.8
Tongue, canned:	Average Maximum	0.8929	0.8565 0.8726 0.8363	58.0	1.4628 1.4677 1.4573	35.8 41.4 26.5	26.7. 33.0 15.0	48.8 63.6 37.8	193 205 181	0.97 2.60 0.42	90.91 94.78 86.20	37.8 46.5 32.4
Fowl:	Average Maximum	0.8999 0.9157 0.8849	0.8633 0.8785 0.8490	56.7 62.5 49.0	1.4637 1.4674 1.4590	31.2 34.0 28.0	19.6 36.5 12.0	76.1 86.4 67.0	198 199 196	0.51 9.92 0.15	95.58 96.03 95.39	45.4 52.0 38.9
Sausage, canned:	Average Maximum Minimum	0.8953	0.8596 0.8850 0.8482	54.6 60.3 49.2	1.4620 1.4660 1.4559	35.2 41.5 27.6	22.3 33.5 12.0	55.0 63.8 39.8	193	0.70	90.76 92.68 88.25	40.04

+ U. S. Dept. of Agric., Bur. of Chem., Bul. 13, part 10, p. 1490 et seq.

accurate, considering the wide variation in nitrogenous content of the various compounds present, a fairly close approximation to the total nitrogenous substance present is obtained by using this factor, since the proteins form by far the largest group of all. The factors to be employed in the calculation of the meat bases are given elsewhere (page 252).

Separation and Examination of Nitrogenous Bodies.—Just how far the analyst should subdivide the various nitrogenous bodies present in meat depends largely on the nature of the case in hand. Frequently the simple determination of total nitrogen as above is sufficient. It is rarely necessary to go further than to divide the nitrogenous bodies into several main groups, according to their solubility in water or other solvents, and their behavior toward certain reagents.

The nitrogen may be determined separately in each of these classes, and by the approximate factor the corresponding nitrogen substance, or class of substances ascertained.

In order to separate most completely the various classes of nitrogenous bodies found in meat, a portion of the fat-free sample should first be exhausted with cold water, which removes the soluble proteins (soluble globulins, proteoses, and peptones) and meat bases, leaving behind the insoluble globulins, the sarcolemma, the albuminoids of the connective tissue (elastin, etc., also insoluble) and the collagen. By next exhausting with boiling water the collagen is removed in the form of soluble gelatin.

By treatment of the combined aqueous extract with zinc sulphate, and with sodium chloride and tannic acid, as hereafter explained, the soluble proteins, including the peptones and gelatin, may be separated from the meat bases.

In obtaining the results from which the table on page 222 was compiled, but three divisions of nitrogenous substances were made, viz., (1) those insoluble in hot water; (2) those precipitated from the water extract by bromine; and (3) the flesh bases. Owing to the incompleteness of the bromine precipitate, the figures given there for nitrogen precipitated by bromine are somewhat high, and those for nitrogen as meat bases are correspondingly low. This fact was observed during the progress of the work, and pointed out in the text with the statement that "considering the small amount of these bodies contained in meat, the results are believed to be approximately correct." See also page 250.

Determination of Nitrogenous Substances Insoluble in Water.—The sample is thoroughly extracted with cold water, the filter and insoluble material transferred to a flask, and nitrogen determined by the Gunning

or Kjeldahl method. The insoluble nitrogen thus obtained is multiplied by 6.25 to obtain insoluble proteins. It is obvious that the insoluble nitrogen may be obtained by difference, the cold water extract being diluted to definite volume, the nitrogen determined in an aliquot portion, and calculated to percentage of soluble nitrogen in the weight of original sample corresponding to the aliquot portion taken. The figure thus obtained, deducted from the percentage of total nitrogen, gives the percentage of insoluble nitrogen.

Trowbridge and Grindley* digest the sample (thoroughly ground in a meat chopper) for one hour in ice water, in the proportion of 1000 grams of meat to 1500 cc. of water. The resulting solution is filtered through cheese cloth, the process being assisted by squeezing the cloth and its contents with the hand. The residue is divided into smaller portions, placed in beakers, and washed in series, using fresh water with No. 1 only, and filtering through cheese cloth from one beaker to another until the last filtrate is colorless, neutral to phenolphthalein, and gives no reaction for proteins by the biuret test. The mixed filtrates and washings filter through paper readily, and give a clear red filtrate.

Pennington† proceeds as follows with the meat of chickens:

A portion of the finely divided red or white meat, weighing 60 grams, is put into a tall, slender bottle of 500 cc. capacity, constructed to fit a centrifuge capable of carrying 1 liter of material; 300 cc. of water are added, and the flask gently shaken for 15 minutes. The movement is merely sufficient to keep the particles of meat in motion and the composition of the extract homogeneous. Forcible shaking causes an emulsion to form, as does the very fine grinding of the tissue. After shaking for the required length of time, the flask is rotated in an electric centrifuge for 20 minutes, which causes the heavier particles to settle in a compact mass, and permits the decantation of the supernatant liquid, which is then filtered through paper. The extraction, as outlined, is repeated with portions of 300 cc. of water until the filtrate is practically protein free, as indicated by the biuret reaction. The attainment of this result requires ordinarily a volume of 1500 to 2500 cc. To guard against bacterial decomposition, thymol is added both to the flesh and to the extract, and to inhibit, so far as possible, the action of the naturally occuring enzymes of the meat, the solution and the meat itself are kept cold, ice being used when necessary.

^{*} Jour. Am. Chem. Soc., 28, 1906, p. 472.

[†] U. S. Dept. of Agric., Bur. of Chem., Bul. 115, p. 64.

The extraction of the white meat is a much simpler operation than the extraction of the dark meat. The latter does not settle as compactly in centrifuging, filters more slowly, and persists in showing a distinct biuret reaction for a considerable time after the white meat is free of water-soluble proteins. In fact, certain fowls, more especially those which have been in cold storage for long periods of time, never show a dark meat entirely free from water-soluble nitrogen. In such cases, the question of the error due to long manipulation and enzyme action, involving a rise in the actual quantity, has to be considered. It has been found by experiment that after long extraction of such tissue, a point is reached when a very faint biuret reaction, which does not apparently diminish, persists indefinitely. Such extractions are halted after about 26 hours, it being believed that a greater error would result in the gain of what has been originally insoluble material, than the loss of the preformed water-soluble nitrogen. The total extract of the muscle is made up to a definite volume, and neutralized to litmus paper with tenth-normal sodium hydroxide.

Cook weighs 200 grams in a 450 Erlenmeyer flask, adds 250 cc. of water, and shakes for three hours in a shaking machine. The material is then filtered by means of linen bags, and extracted with water repeatedly by vigorous manipulation with the hands in successive portions of water, pressing out after each extraction until negative biuret reaction is obtained. The operation ordinarily requires from 2200 to 2500 cc. of water. A small quantity of phenol and thymol are added as preservatives.

Weber* applied Cook's method at room temperature and with ice water to samples of fresh and storage meat, as well as to samples which he had kept for varying lengths of time in the laboratory. He obtained a larger amount of soluble proteins when working at room temperature. No opinion was expressed as to whether this was due to the greater extracting power of water at room temperature, or to greater enzymic action during the period of extraction.

Determination of Collagen.—The insoluble proteins obtained as directed above are transferred to a beaker, water added, and heated to boiling for some minutes. They are then separated by filtration and washed with boiling water. The nitrogen of the residue insoluble in boiling water is deducted from the nitrogen insoluble in cold water, and multiplied by 5.55 for the per cent of collagen. This method is of doubt-

^{*} U. S. Dept. of Agric., Bur. of Chem., Bul. 122, p. 42.

ful value, owing to the difficulty of converting the collagen to soluble form on the one hand, and to the tendency to decompose a portion of the protein on the other.

Determination of Coagulable Proteins.—The entire filtrate, or an aliquot portion thereof, from the determination of nitrogenous bodies insoluble in water is heated sufficiently to coagulate the coagulable proteins, filtered, the insoluble material washed with hot water, and the filter and contents transferred to a Kjeldahl flask and nitrogen determined by the Gunning method. The per cent of nitrogen multiplied by 6.25 gives the per cent of coagulable proteins.

The amount of heating necessary to obtain maximum coagulation varies with different materials. The Association of Official Agricultural Chemists directs that the solution be almost neutralized, but left still faintly acid, and boiled until the globulins are coagulated.*

Pennington,† working with chickens, evaporates 350 cc. to a volume of about 100 cc. before filtering. Grindley and Emmett‡ employ 200 cc. of the solution, add alkali till neutral to litmus paper, and evaporate to 50 cc. In a later article, Trowbridge and Grindley § report maximum results from the cold water extract of fresh beef by neutralizing one-fourth of the acidity to phenolphthalein before coagulation.

Determination of Proteoses, Peptones, and Meat Bases.—The filtrate from coagulated proteins, having been diluted by wash water, is concentrated by evaporation and made up to 100 cc. Proteoses are then determined by Bömer's method (page 250), and meat bases by Sjerning's method, as modified by Bigelow and Cook (page 252). Peptones are determined by difference—the sum of the nitrogen occurring in insoluble nitrogenous bodies, coagulated proteins, meat bases and ammonia being deducted from the total nitrogen.

Determination of Gelatin.—Modified Stutzer's Method. —A weighed portion of the sample, say 10 grams, is thoroughly extracted by boiling water, the extract transferred to a porcelain dish containing about 20 grams of previously ignited sand, and evaporated to dryness. The residue is then stirred with four successive portions of absolute alcohol,

^{*} U. S. Dept. of Agric., Bur. of Chem., Bul. 107 (rev.), p. 108.

[†] Ibid., Bul. 115. p. 65.

[‡] Jour. Am. Chem. Soc., 27, 1905, p. 665.

[§] Ibid, 28, 1906, p. 494.

^{||} U. S. Dept. of Agric., Bul. 13, part 10, p. 1307.

using about 50 cc. each time and pouring it off through a filter consisting of a layer of asbestos fiber on a perforated porcelain plate within a funnel. This funnel is surrounded by chopped ice, and is so arranged that gentle suction may be used to hasten the filtration. The residue is then repeatedly stirred with successive portions of about 100 cc. each of a mixture containing 100 cc. of 95% alcohol, 300 grams of ice, and 600 grams of cold water, the portions being passed through the asbestos filter, and the washing being continued till the solution is colorless as it comes from the filter, keeping the temperature always below 5° . The asbestos is then transferred to a beaker with the washed residue, and the whole thoroughly extracted with boiling water. The hot-water extract is evaporated to small volume, and washed into a Kjeldahl flask, in which it is then evaporated to dryness, and the nitrogen determined by the Gunning method: $N \times 5.55 = \text{gelatin}$.

Detection of Nitrates.—A small portion of the finely divided material is treated in a porcelain dish or on a tile with a little 1% solution of diphenylamine in concentrated sulphuric acid. Presence of a blue color indicates nitrate.

Detection of Preservatives.—Meats may be systematically tested for preservatives in the same manner as canned goods. The preservatives most commonly used in meat and meat preparations are tested for as follows:

Detection of Sulphurous Acid.—Proceed as directed on page 834.

Traces should be ignored, as slight reactions for sulphurous acid are obtained with meats that have not been chemically preserved. This is probably due to the decomposition of a portion of the proteins. According to Mentzel,* 4 milligrams per 100 grams may be due to this cause.

Detection of Boric Acid.—A portion of the finely divided meat, mechanically freed from the fat as far as possible, is warmed with water acidified with hydrochloric acid, and turmeric-paper is soaked in the extract. The rose-red color of the turmeric-paper after drying (turned blue by weak alkali) is indicative of boric acid.

A more delicate method of procedure consists in burning to an ash a portion of the meat, after treatment with lime water, and testing with turmeric tincture a solution of the ash slightly acidified with hydrochloric acid.

^{*} Zeits. Unters. Nahr. Genuss., 11, 1906, p. 320.

Detection of Salicylic Acid.—The sample, mechanically freed from fat, is slightly acidified and shaken out with ether. The ether extract evaporated to dryness is tested with a drop of a solution of ferric chloride. A deep-violet coloration indicates salicylic acid.

Starch in Sausages, Meat-balls, etc.—Detection.—The addition of cracker or bread crumbs is best indicated by the presence of considerable starch, which is readily recognized by the iodine test, applied by boiling up a portion of the sample with water, cooling and adding a drop of iodine reagent to the liquid. The characteristic blue color is produced, if starch be present in notable quantity. Traces of starch may be due to the pepper and spices used in seasoning the sausage. A small admixture of starch is rendered apparent when a thin section of the sausage is treated with a drop of iodine reagent and viewed under the microscope. A microscopical examination will sometimes reveal the character of the starch, whether it is from cereals or from pepper, but in some preparations the starch is thoroughly cooked and its structure destroyed.

Estimation of Starch.—The regular acid conversion process, p. 283, may be applied, but more accurate results are obtained by the method of inversion with malt extract. Medicus and Schwab* prepare the malt extract for this purpose by digesting 5 grams of ground malt with 50 cc. of water for one and one-half hours at 20° to 30° C. In making the starch estimation, they digest for two hours at a temperature of from 40° to 50° C. 20 grams of the sausage mixed with 20 cc. of the malt extract, and afterwards for eighteen hours at room temperature. After filtering and washing, the filtrate is boiled to coagulate the albumin and again filtered. The second filtrate is then made up to 200 cc., 20 cc. of 25% hydrochloric acid (specific gravity 1.125) are added, and the starch determined in the regular manner.

Mayrhojer's Method.†—This is considered the simplest and most reliable method of estimating the starch in such substances as sausages. From 60 to 80 grams of the sample are heated on the water-bath with an 8% solution of alcoholic potassium hydroxide, which, in the case of pure sausages, dissolves nearly everything except a little cellulose. To prevent gelatinization, warm alcohol is added to dilute the solution, which is then filtered through paper or asbestos. The starch is con-

^{*} Berichte d. chem. Gesell., XII, p. 1285.

[†] Zeits. Nahr. Untersuch., 1896, p. 331; Abs. Analyst, 1897, p. 11.

tained in the insoluble residue, which is washed with alcohol till the washings are no longer alkaline, after which it is treated with an aqueous solution of potassium hydroxide, and the starch solution made up to a definite volume. To an aliquot part of the solution 95% alcohol is added, whereupon the starch comes down as a flocculent precipitate. This is collected on a weighed filter, washed with alcohol and ether, dried, and weighed. The filter with its contents is then burnt to an ash, the amount of which is deducted.

In order to avoid the ash determination, the starch may be precipitated from a weak acetic acid solution instead of from an alkaline solution, the potassium acetate formed being soluble in the alcohol, and nothing but pure starch is precipitated.

Characteristics of Horse Flesh.—Although certain authorities have found distinguishing characteristics in color, consistency, odor, etc., between horse flesh on the one hand, and beef and pork on the other, it is extremely difficult, by its physical properties, to detect horse flesh when mixed with other meat, especially when the mixture is chopped. Horse flesh has a much coarser texture and is darker in color than beef. The muscle fibers are, as a rule, shorter in horse flesh. On treating horse flesh with formal-dehyde, Ehrlich* has found that a very characteristic odor is developed within forty-eight hours, suggestive of roasted goose flesh.

Certain of the constants of the fat of horse meat differ from those of beef and pork, notably the iodine value and the refractometer readings. These constants are compared as follows:

	Iodine Value.	Butyro-refractom- eter Readings. Temperature 40°.
Horse fat	71–86 38–46 50–70	53-7 49-0 48-6-51-2

The fact that glycogen usually exists to a much larger extent in horseflesh than in other meat, renders it possible in some cases to detect horseflesh, when present in the mixture.

The following table prepared by Bujard shows the relative amount of glycogen in various kinds of meat and sausages:

^{*} Zeit. Fleisch u. Milch Hyg., 1895, p. 232.

	Water.	Glycoge	n Direct.	Glycogen in Dried Substance.		
		Niebel Method.	Mayrhofer Method.	Niebel.	Mayrhofer.	
Horse flesh	74-44 74-87	0.440	0.445	1.721	1.741	
"	76.17 76.00	1.827 0.592	0.610	7.667 2.466	7.247	
Red sausage (Knackwurst) Pork sausage Veal.	69.26 67.25 74.6		0.038		0.124 0.733 0.342	
Pork.	75.0		0.186	•••••	0.744	

In beef Bujard found 0.74 and 0.073 per cent of glycogen calculated in terms of dried substance, and, in sausages made exclusively from horse meat, amounts of glycogen ranging from 0.05 to 5.34, the sample in the latter case being made from the liver. It was formerly thought possible to detect as small an amount as 5% of horse flesh in mixture, but later investigation showed that after the death of the animal, glycogen, though present at first in considerable quantity, decomposes more or less rapidly, going over into muscle sugar (dextrose). Hence, while the presence of much glycogen is suspicious, its absence is by no means proof that horse flesh was not used.

Niebel did not consider the failure of the glycogen test as sufficiently conclusive to establish the absence of horse flesh, on account of the tendency toward decomposition of the glycogen. In the absence of starch, he regards the presence of more than 1% of dextrose in the fat-free meat, after conversion of the carbohydrates, to be proof of the presence of horse-flesh.

Detection of Glycogen.—From the well-known color reaction produced by iodine on glycogen, horse flesh can often be detected, when present in sausages, unless obscured by the presence of starch or dextrin.

Brautigam and Edelmann * proceed as follows: 50 grams of the finely divided meat are boiled with 200 cc. of water for an hour, and, after cooling, dilute nitric acid is added to the broth to precipitate the proteins and to decolorize. The broth is then filtered, and a portion of the filtrate is treated in a test-tube with a freshly prepared, saturated, aqueous solution

^{*} Pharm. Central., 1898, p. 557.

of iodine, or, better, with a mixture of 2 parts iodine to 4 parts potassium iodide and 100 parts water, the reagent being carefully added so as not to mix with the broth, but form a layer above it. If glycogen be present in considerable amount, a wine-colored ring is observable at the junction of the two layers. On heating the test-tube, the coloration disappears if due to glycogen, but it reappears on cooling. This reaction was found to occur with horse flesh and not with beef, mutton, veal, or pork.*

If the color is not clearly apparent, the chopped meat is heated on the water-bath with a solution of potassium hydroxide (using an amount of potassium hydroxide equivalent to 3% of the weight of the flesh) till the fiber is decomposed, after which the broth is concentrated to half its volume, treated with nitric acid to precipitate the proteins, filtered, and treated with the iodine solution as previously.

Determination of Glycogen.†—Niebel's Modification of Brücke's Method.—This method is applicable only in the absence of dextrose and dextrin. If therefore the presence and character of the starch indicates the presence to a considerable extent of cracker crumbs or other cereal "filler," the method is not accurate.

A weighed portion of the flesh is heated on the water-bath with 3 to 4 per cent of potassium hydroxide and four volumes of water for six hours. Evaporate the broth to half its original bulk, and add, after cooling, a solution of mercuric iodide in potassium iodide,‡ which precipitates the protein. Filter, and to the clear filtrate add $2\frac{1}{2}$ times its volume of 95% alcohol, collect the precipitated glycogen on a filter, wash first with 60% alcohol, then with 95% alcohol, then with absolute alcohol, then with ether, and finally with absolute alcohol. Dry at 115° C. and weigh.

Landwehr's Method.—Applicable in presence of dextrose. The broth prepared as in Niebel's method is freed from protein by the addition of zinc acetate. Filter, wash, and heat the entire filtrate on the waterbath with sufficient of a concentrated solution of ferric chloride, afterwards precipitating the iron with a few drops of a saturated solution of sodium hydroxide. Filter, wash the precipitate with hot water, and dis-

^{*} The reaction was found to occur also with the flesh of the human foctus and with the foctus of animals; also with mule meat, but not with the flesh of the dog or cat.

[†] Jahresb. Nahr. Genuss., 1891, p. 38.

[‡] The reagent known as Brücke's reagent is prepared by precipitating a solution of mercuric chloride with potassium iodide, washing the precipitated mercuric iodide till free from chloride, and afterwards saturating, while boiling, a 10% potassium iodide solution with the mercuric iodide.

solve in strong acetic acid. Add to the solution, after cooling, sufficient hydrochloric acid to produce a yellow color, then pour into 2½ volumes of alcohol, and proceed as in the preceding paragraph.

Mayrhofer's Method,* on which the results in the table on page 235 are based, is as follows: Dissolve a weighed portion of the flesh in an aqueous solution of potassium hydroxide, precipitate the proteins by hydrochloric acid and Nessler's reagent, filter, and treat the clear filtrate with alcohol, which precipitates the glycogen. This is collected on a tared filter and washed, first with dilute alcohol, and finally with ether, dried at 110° C., and weighed.

Pflüger and Nerking's Method.†—Of the finely divided sample 50 grams are heated on the water-bath with 200 cc. of 2% potassium hydroxide till the solution is practically complete. After cooling, the solution is made up to 200 cc. with water, shaken, and filtered. To 100 cc. of the filtrate, 10 grams of potassium iodide and 1 gram potassium hydroxide are added, and the solution stirrred till clear, after which 50 cc. of 95% alcohol are added and the mixture allowed to stand over night. precipitates the glycogen. Filter, and wash the precipitate with a solution made up of 1 cc. 70% potassium hydroxide, 10 grams potassium iodide, 100 cc. water, and 50 cc. 95% alcohol. After further washing the glycogen with 2 parts strong alcohol and 1 part water, dissolve in water and by means of Brücke's mercuric-iodide-in-potassium-iodide reagent (see footnote, p. 236) remove any remaining nitrogenous substances. turbid, and to the solution add common salt (about 2 milligrams per 100 cc. of solution), and reprecipitate the glycogen by adding 2 volumes of 95% alcohol. Filter, wash first with 95% alcohol containing a little common salt, then with absolute alcohol, and lastly with ether. Dry and weigh.

Bigelow suggests that the glycogen as above obtained be converted by acid hydrolysis to dextrose, which is determined in the regular manner. Dextrose \times .0=glycogen.

Identification of Raw Horse Flesh by the Blood Serum Test.‡—This test depends upon the recent development of the principle that when a rabbit has been inoculated with the blood of a particular animal, as for instance that of the horse, the serum of the rabbit's blood will react with

^{*} Forsch. Ber., 1897, IV, 47.

[†] Arch. ges. Physiol., 1899, 76, 531-542; Bul. 65, Bur. of Chem., p. 13. Recommended for Provis. Adoption by the A. O. A. C.

^{\$} Schaltze, A., Ueber weitere Anwendungen der Präcipitine. (Deuts. med. Wochs., 1902, No. 45, p. 804.)

Wassermann, A., u. Schütze, A., Ueber die Entwickelung der biologischen Methode

the blood of the horse and with that of no other animal. To prepare the blood serum for a reagent, inject a rabbit with 10 cc. of defibrinated horse's blood every day for five to six days, either subcutaneously or intravenously. The blood afterwards taken from the rabbit is clotted, and the filtered serum is used in making the test, or, if the reagent is to be kept for some time, the rabbit's blood serum is dried and an aqueous solution used for the reagent.

If the clear expressed juice from the suspected flesh, filtered if necessary, be treated with a few drops of the rabbit's blood reagent, prepared as above, a cloudy precipitate will be produced in the case of horse flesh.

By inoculating different rabbits in like manner with the blood of various animals, the flesh of the corresponding animals may be recognized from the reaction of the blood serum of the rabbit with its juices. Only raw flesh responds to the test, as heating destroys the virtue of the reagent.

Determination of Muscle Sugar (Dextrose).—Boil a weighed quantity of the finely divided sample, say 50 grams, with water, add an excess of normal lead acetate solution, and make up with water to a given volume, say 250 cc. Filter, and to an aliquot part of the filtrate add enough of a saturated solution of sodium sulphate to precipitate the lead. Again filter, make up to a given volume, and determine the dextrose in a measured part of the solution by either of the regular methods.

Detection of Coloring Matter.—Red Ocher is indicated by an excessive amount of iron in the ash.

Cochineal is most readily tested for by the method of Klinger and Bujard.* The sausage, finely divided, is heated with two volumes of a mixture of equal parts of glycerin and water for several hours on the water-bath, the mixture being slightly acidified. The yellow solution is passed through a wet filter, and the coloring matter, if present, is precipitated as a lake by adding alum and ammonia, the precipitate is filtered off and washed, after which it is dissolved in a small amount of tartaric acid, and the concentrated solution, contained in a test-tube, is examined

zur Unterscheidung von menschlichem und tierischem Eiweissmittels Präcipitine. (Ibid., 1902, No. 27, p. 483.)

Wassermann, A., Ueber Agglutinine und Präcipitine. (Zeits. f. Hyg., etc., Bd. 42, 1903, 2, p. 267.)

Uhlenhuth, Die Unterscheidung des Fleisches verschiedener Tiere mit Hilfe spezifische Sera und die praktische Anwendung der Methode in der Fleischbeschau. (Deuts. Med. Wochs., 1901, No. 45, p. 780.)

Miessner, H., u. Herbst, Die Serum agglutination und ihre Bedeutung für die Fleis huntersuchung. (Arch. f. wissensch. u. prakt. Tierheilk., 1902, Heft 3-4, p. 359.)

^{*} Zeit. angew. Chem., 1891, p. 515.

through the spectroscope for the characteristic absorption-bands of carmine lake, lying between b and D.

Spaeth * has shown that both carmine (cochineal) and anilin red, which are the dyes most commonly used for coloring sausages, can be most readily extracted therefrom by warming the finely divided material a short time on the water-bath with a 5% solution of sodium salicylate.

Vegetable and Coal-tar Colors.—Various solvents are suggested for the removal of these dyes from sausage meat. Allen† recommends extraction with methylated spirit (a mixture of ethyl alcohol with 10% methyl); Bigelow‡ recommends heating with a mixture of 50% glycerin slightly acidified; A. S. Mitchell uses alcohol acidified with hydrochloric acid; Spaeth a 5% solution of salicylate of soda. Other solvents applicable in some cases are dilute ammonia and amyl alcohol. The solvent, after filtering, is evaporated to small volume, acidified with hydrochloric acid, and white wool is boiled in it. If the wool is distinctly dyed, a coal-tar color is undoubtedly present, and this can often be identified by methods given in Chapter XVII. According to Marpmann, pure normal flesh containing natural color only is completely decolorized by macerating for two hours in 50% alcohol, while artificially colored meat remains colored after this treatment.

Marpmann's Microscopical Method. —Moisten a thin section of the sausage with 50% alcohol, and examine under the microscope. Some colors are readily apparent without further treatment. If only traces of color are present, clarify the substance by treatment with xylol, which is removed by the use of carbon tetrachloride. The mass rendered transparent by this treatment is then immersed in cedar oil and examined, the coloring matters, if present, being especially apparent. If the color used is fuchsin (magenta), carmine, logwood, or orchil, the substance of the cell will appear stained. If acid coal-tar dyes are used, the liquid contents of the cell will show the color.

Detection of Frozen Meat.—Maljean || detects frozen meat by the aid of a microscope. A drop of the blood or meat juice is pressed out upon a slide and immediately examined before it solidifies. Fresh meat juice contains many red blood corpuscles, floating in a clear colorless serum, and readily apparent. In blood from frozen meat, the red cor-

^{*} Pharm. Central., 1897, 38, p. 884.

[†] Commercial Organic Analysis, Vol. IV, p. 294.

[†] U. S. Dept. of Agric., Bureau of Chemistry, Bul. 65, p. 16.

Zeits. angewand. Mikrosk, 1895, p. 12.

[|] Jour. Pharm. et Chem., 1892, XXV, p. 348.

puscles are nearly always completely dissolved in the serum, due to freezing, or, if not dissolved, are much distorted and entirely decolorized, the liquid portion being darker than usual.

Megascopically, the fresh meat juice is more abundant than that of frozen meat, and its color is deeper. According to C. A. Mitchell, if a small piece of meat once frozen be shaken in a test-tube with water, color is imparted to the water much more quickly than with fresh meat, and the color is deeper.

MEAT EXTRACTS.

Character and Composition.—Numerous preparations sold under the name of meat extracts have been on the market for many years. At the beginning of the nineteenth century the value of such extracts was known, but Liebig was the first some fifty years later to produce a commercial extract of meat. Liebig's preparation, as originally made, consisted of a cold-water extract of chopped lean meat, strained free from fiber, heated, filtered, and evaporated, thus containing little of any gelatin or proteins. Later, however, Liebig advocated the use of warm and even boiling water for extraction, by which method of preparation the amount of gelatin was greatly increased. He, however, condemned the use of salt.

The best modern meat extracts consist for the most part of such portions of the meat, previously freed from bone and superfluous fat, as are soluble in water the temperature of which does not exceed 75° C. The widest latitude, however, prevails as to the temperature employed for the extraction, hence the character of the various products is somewhat varied. It is not an uncommon practice to submit the meat to actual boiling with water, in which case the amount of gelatin will be considerable. In an extract prepared by warm water, one finds very little gelatin, more or less albumin, albumoses, and peptones, and practically all the meat bases, phosphates, and chlorides present in the meat: also minute quantities of lactic acid, inosite, and possibly glycogen. By far the most important of these substances from the physiological standpoint are the meat bases—creatin, creatinin, xanthin, sarkin, etc. To the predominance of these amido-bodies is undoubtedly due the wellknown stimulating effect of meat extracts. Indeed, a properly prepared extract has very little actual food value, but is rather to be regarded as a condiment or as a stimulant, acting on the nervous system in a somewhat analogous manner to tea and coffee.

Commercial meat extracts differ much in consistency according to

the extent to which evaporation is carried, varying from the thin fluid through the pasty form to the semi-solid. Some preparations have added thereto finely ground dried beef or beef meal.

Owing partly to unfounded claims of manufacturers, meat extracts are commonly confused with meat juices, and products belonging to the former class are sold for the latter. Considering the widely different nature of meat extracts and meat juices, such confusion is a serious matter. Meat extract is employed as a stimulant in the form of beef tea or as a flavor for soups. Meat juices, on the other hand, are employed in the sickroom as a readily available form of food.

Standards.—The following standards for products of this class have been adopted by the Association of Official Agricultural Chemists and the Association of State and National Dairy and Food Departments:

- 1. Meat extract is the product obtained by extracting fresh meat with boiling water, and concentrating the liquid portion by evaporation after the removal of fat, and contains not less than 75% of total solids, of which not over 27% is ash, and not over 12% is sodium chloride (calculated from the total chlorine present), not over 0.6% is fat, and not less than 8% is nitrogen. The nitrogenous compounds contain not less than 40% of meat bases, and not less than 10% of creatin and creatinin.
- 2. Fluid meat extract is identical with meat extract, except that it is concentrated to a lower degree, and contains not more than 75, and not less than 50% of total solids.
- 3. Bone extract is the product obtained by extracting fresh trimmed bones with boiling water and concentrating the liquid portion by evaporation after removal of fat, and contains not less than 75% of total solids.
- 4. Fluid bone extract is identical with bone extract, except that it is concentrated to a lower degree and contains not more than 75 and not less than 50% of total solids.
- 5. Meat juice is the fluid portion of muscle fiber, obtained by pressure or otherwise, and may be concentrated by evaporation at a temperature below the coagulating point of the soluble proteins. The solids contain not more than 15% of ash, not more than 2.5% of sodium chloride (calculated from the total chlorine present), not more than 4 nor less than 2% of phosphoric acid (P_2O_5) , and not less than 12% of nitrogen. The nitrogenous bodies contain not less than 35% of coagulable proteins, and not more than 40% of meat bases.
 - 6. Peptones are products prepared by the digestion of protein material

ANALYSES OF SOLID MEAT EXTRACTS.

	Undetermined.	8.66 5.89 11.67 16.11 21.04
	Ether Extract.	1.30 0.94 0.50 0.53 0.43
.ainommA		0.20 0.37 0.43 0.71
	Meat Bases Other than Creatin and Xanthin.	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2
	Xanthin Bases.	0.38 0.03 0.04 0.11 0.45
,	Creatin and Creatinin.	0.87 1.14 0.75 1.01 0.81
Nitrogen as-	Total Meat Bases.	3.56 3.82 3.05 3.20 4.21
Nitrog	Peptones.	1.57 2.68 1.90 1.93 1.03
_	Proteoses.	1.65 2.01 2.02 0.77 1.02 0.86
	Insoluble and Coagulable.	0 . 3 2 0 0 . 3 2 0 0 . 3 5 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
	Total Nitrogen.	7.30 9.07 7.66 6.02 6.60
lity.	As Lactic Acid.	6.01 88.13 8.42 5.15 6.44
Acidity	N/10 Sodium Hydroxide, cc. per gram.	6.67 9.04 9.35 5.72 4.61 7.16
	Inorganic Phos- phoric Acid.	1.94 1.79 2.29 2.71 2.98
tuents	Organic Phos- phoric Acid.	0.35 0.61 0.24 0.18 0.18
Consti	Total Phosphoric Acid.	2 . 2 0 2 . 4 0 2 . 5 3 2 . 5 3 3 . 1 9
Mineral Constituents.	Chlorine as So- dium Chloride in Ash.	8.54 3.11 5.47 18.32 13.51
4	.ńsA latoT	24.06 21.03 20.46 30.92 31.68
	Moisture.	26.50 21.14 21.66 21.86 20.16
		Cudahy's Rex brand. Liebig's Armour's Libby, McNeill & Libby's Premier Swiff's Hammond's Coin Special

NITROGENOUS CONSTITUENTS OF SOLID MEAT EXTRACTS (CALCULATED PROM ABOVE).

	trogen.	sinommA	2.74 4.08 2.74 7.14 10.76 3.64
	Nitrogenous Bodies Expressed in Terms of Total Nitrogen.	Meat Bases Other than Creatin, Creatinin, and Xanthin.	31.64 29.32 29.50 34.55 32.88
.	rms of	Xanthin Bases.	7.00.33
	in Te	Creatin and Creatinin.	11.92 12.57 9.79 16.78 12.27 18.08
	Kpresse	Total Meat Bases.	48.77 42.12 39.82 53.16 51.97 61.37
	odies E	Peptones.	21.51 29.55 24.80 22.09 16.52
	enous B	Proteoses.	22.60 22.16 26.37 12.79 15.45
	Nitroge	Insoluble and Coagulable Protein.	8.4.0 6.0.0 7.88.4 7.80.0 7.80.0
		.sinommA	2.000 4.4.000 6.0000 6.0000
		Meat Bases Other than Creatin, Creatinin, and Xanthin.	7.21 7.05 6.49 6.77 7.64
		Xanthin Bases.	0.08
	dies.	Creatin and Creatinin.	3.36 3.36 3.34 3.53 3.63
	ous Bo	Total Meat Bases.	11.11 11.92 9.52 9.98 10.70
	Nitrogeneous Bodies.	Peptones.	9.81 11.888 11.883 6.81 9.25
	Z	Proteoses.	12.50 12.50 12.50 6.38 5.38
		Insoluble and Coagulable Pro- tein.	2.00 1.19 3.00 1.81 2.19
		*.anistor¶ latoT	22.12 30.50 27.51 14.93 15.38
			Cudahy's Rex brand Liebig's Limout's Libby, McNeill & Libby's Premier Swift's Hammond's Coin Special

* The sum of insoluble and coagulable proteins, proteoses, and peptones.

EXTRACTS.
MEAT
FLUID
OF
ANALYSES

i		Undetermined.	9.00 12.12 12.00 13.54 13.54		
		Ether Extract.	0.00000		
		.sinommA	0000000		
		Meat Bases other than Creatinin and Xanthin.	1.05		
		Xanthin Bases.	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0		
	ļ	Creatin and Creatinin.	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0		
	Nitrogen as-	Total Meat Bases.	1.06 1.96 1.06 2.06 1.36 1.36	ble.	
	Nit	Peptones.	0.70 0.74 0.07 14.00 0.80 19.00	t Partly insoluble.	
		Proteoses.	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	Partly	
2010		bns sidulosen Cosgulable Protein.	0.04 0.04 0.03 0.03 0.03 0.03	#	
- 1		Total Mitrogen.	3.15 3.06 3.06 3.87 3.98 3.18		
THE COLD	Acidity.	As Lactic Acid.	3.11 3.92 4.53 4.92 2.43 2.20	luble.	
	Acic	N/10 Sodium Hydroxide, cc. per gram.	3.46 4.35 5.04 5.29 5.31 2.70	Largely insoluble.	
		Inorganic Phos- phoric Acid.	2.06 2.90 2.83 2.10 0.81 0.62	Large	
	tuents.	Organic Phos- phoric Acid.	0.20 4.00 4.00 6.13 8.10 8.10	+	
TO GET THE	Consti	Total Phosphoric Acid.	2 8 8 8 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9		
	Mineral Constituents.	Chlorine as So- dium Chloride in Ash.	8.27 6.71 7.02 7.02 8.48 11.38	ulable	
	*	Total Ash.	17.23 16.21 10.26 15.91 16.99 16.13	All coagulable.	
		Moisture.	57.75 58.84 57.64 49.94 55.99 64.63	7.	
İ		•	Beef Ext.		
			rmour's Conc Pluid Beef Ext. yeth's Beef Juice alentine's Meat Juice rmour's Vigoral dahy's Rex Pluid Beef Ext. bil's Pluid Beef Ext osquera Pluid Beef Juic		
			Conc Pluid Bee Jeef Juice s Meat Juice Vigoral Ker Pluid Beef Ext. Pluid Beef Ext. Pluid Beef Jell.		
			our's Concina Beef this Beef trine's Mour's Vige hy's Rex s Fluid Beef uers Fluid B		
			Armour's Wyeth's Valentine Armour's Cudahy's Cibil's Flu		

NITROGENOUS CONSTITUENTS OF FLUID MEAT EXTRACTS (CALCULATED FROM ABOVE).

trogen.	.ainommA	5.35 5.35 5.35 5.35 5.35 5.35
Total Ni	Meat Bases other than Creatin, Creatinin, and Xanthin.	36.84 44.44 44.47 35.40 45.32 24.21 26.56
rms of	Xanthin Bases.	3.32 3.32 3.32 3.32
d in Te	Creatin and Creatinin.	13.33 11.44 12.45 10.72 10.72
Nitrogenous Bodies Expressed in Terms of Total Nitrogen	Total Meat Bases.	58.25 63.95 63.40 66.58 42.77
odies E	Peptones.	24.56 25.16 19.12 10.38 27.99
mous B	Proteoses.	11.93 3.17 3.27 17.83 13.67 13.84 12.86
Nitroge	Insoluble and Coagulable Protein.	1.40 0.98 0.98 7.49 9.75
	Ammonia.	000000
	Meat Bases other than Creatin, Creatinin, and Xanthin.	8 4 4 4 8 8 8 8 9 9 9 9 9 9 9 9 9 9 9 9
	Xanthin Bases.	0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.0
odies.	Creatin and Creatinin.	1.19 1.09 1.50 1.50 1.50 1.50
neous B	Total Meat Bases.	3.05.99 3.05.30 3.05.30 3.05.30
Nitrogeneous Bodies	Peptones.	8 4 4 4 8 8 8 4 4 6 9 4 8 8 8 9 4 8 9 4 8 9 9 9 9 9 9 9 9 9
•	Proteoses.	2.13 0.03 0.03 4.31 2.75 1.94
	Insoluble and Coagulable Pro- tein.	2 . 0 . 0 . 0 . 0 . 0 . 0 . 0 . 0 . 0 .
	*.enistor4 latoT	5.76 5.63 10.75 10.25 8.13
		Armour's Conc. Fluid Beef Ext. Wyeth's Beef Juice Welth Steef Juice Armour's Vigoral Cudahy's Rex Fluid Beef Ext Cibil's Fluid Beef Ext. Mosquera Fluid Beef Ext.

* The sum of insoluble and coagulable proteins, proteoses, and peptones.

by means of enzymes or otherwise, and contain not less than 90% of proteoses and peptones.

7. Gelatin (edible gelatin) is a purified, dried, inodorous product of the hydrolysis, by treatment with boiling water, of certain tissues, as skin, ligaments, and bones, from sound animals, and contains not more than 2% of ash and not less than 15% of nitrogen.

ANALYSES OF MEAT EXTRACTS.—Largely by the application of the above standards, Bigelow and Cook* have classified a number of products of this class as solid (pasty) meat extracts, fluid meat extracts, and "miscellaneous preparations."

Their results are given on pages 242, 243, 247, and 248.

Solid and Fluid Meat Extracts.—It will be noted that the solid and fluid extracts are identical, except that the latter are concentrated only half as much as the former. Allent holds that the maximum chlorine content of meat extract calculated to sodium chloride is 0.06% for every unit of dry solid matter, and that excess over that amount is due to added common salt. This opinion is based on the composition of South American extracts prepared from the meat of the entire carcass. Street! considers that the maximum standard of 12% is too high, and encourages the manufacturer to add salt to his product. In this country, however, extracts are commonly prepared in part by the evaporation of the soup liquor in which meat is parboiled before canning, and in part from trimmings. It is claimed that the natural salt content of the product made in this manner is higher than when the entire meat of the carcass is employed. A second grade article is also made from bones, trimmings, etc., and contains a still higher percentage of sodium chloride. This product is designated as "bone extract" in the standards given on page 241.

The presence of an excessive amount of sodium chloride is usually due, probably, to the presence of the product last mentioned, or to the use of corned beef in the preparation of the substance. In the latter case nitrates are generally present. On comparing the analyses given above with the composition of other products of this class, as contained in the following tables, the value of the percentage of meat bases, especially of creatin and creatinin, in distinguishing meat extracts from meat juices and manufactured products of that general type is apparent.

^{*} U. S. Dept. of Agric., Bur. of Chem., Bul. 114.

[†] Commercial Organic Analysis, vol. 4, p. 307.

^{. ‡} Conn. Expt. Station, Report for 1907 and 1908, p. 622.

[§] Bigelow and Cook, U. S. Dept. of Agric., Bur. of Chem. Bul. 13, pt. 10, p. 1389. Bigelow and Cook, U. S. Dept. of Agric., Bur. of Chem., Bul. 114, p. 13.

Meat Juices Prepared in the Laboratory.—For the purpose of comparison with meat extracts, the following analyses of meat juices prepared in the laboratory are of interest.

MEAT JUICES PREPARED IN LABORATORY.*

		Composition of Sample.								
Preparation of Juice.		Water in Juice	A:	sh.	Chlori as Sodiu Chlori in As	m pho de (Pa	oric eid O ₆).	Ether Extract.	Acidity as Lactic Acid.	
Round beef, cold pressed			85.76 1.53 86.85 1.86 90.65 1.36 91.90 1.29 89.56 1.27		0.2 0.1 0.1	0.12 0.3 0.20 0.3 0.15 0.3 0.19 0.2 0.16 0.3		.31 0.30 .36 0.19 .29 0.64		
Juice pressed from sirloin steal Juice extracted from sirloin s	teak by cold	96.13	1	46	0.0		18			
Juice extracted from beef ch	uck by cold	96.58		43	0.0	-	11			
Juice extracted from beef ch pressure after 6 hours at 60°	uck by cold	98.11	1	39	0.0	1	12			
		1		Co	mpos	ition of	Sample	B.		
Preparation of Juice.			Insolution ble Nitro	o- Ni	ag- able tro- en.	Pro- teose Nitro- gen.	Pep- tone Nitro gen	Amido Nitro- gen.	Unde- ter- mined Matter.	
Round beef, cold pressed		2.08 1.74 1.16 1.09 1.09	0.12	0.16 1.37 0.29 0.98 0.68 0.12 0.41 0.49 0.54		0.06 0.07 0.04 0.07 0.42 0.20	0.16 0.11 0.01 0.21	0.33 0.29 0.43 0.27 0.18 0.26	0.47 1.03 1.90 0.40 2.92 0.94	
pressure Juice extracted from beef chipressure Juice extracted from beef chi	uck by cold	0.48	0.34			trace trace	none none	0.14	0.85	
pressure after 6 hours at 60°-	-100° C'	0.24	0.00			trace	0.12	0.08	0.25	
	Results in	Terms Nitroge		otal		Nit	rogeno	ıs Bodies	i.	
Preparation of Juice.	Insol- uble uble Pro- tein.	Albu- moses		Amide Bodie		le ulabi o- Pro-	e Pro			
Round beef, cold pressed	7.69 65.87 16.66 56.32 58.62 11.01 37.61 44.95	2.88 4.02 3.45 6.42 38.53	7.69 6.32 0.86 19.26	37.0	6 I. 7 7 O.	00 8.56 81 6.13 4.25 75 2.56 3.06	0.4	4 0.69 5 0.06 4 1.31	0.90 1.34 0.84	
and water	45.76	16.95	15.25			3.38	I.2	- 1		
steak by cold pressure Juice extracted from beef chuck by cold pressure Juice extracted from beef	70.83 79.07			20.9	1	2.13	trac	1	1	
chuck by cold pressure after 6 hours at 60°-100° C	0.00	 	50.00	33 - 33	3	0.00	trac	e 0.75	0.25	

^{*} Bigelow and Cook, U. S. Dept. of Agric., Bul. 114, p. 19.

The composition of these products is widely different from that of the so-called meat juices of commerce, as given in the table on page 247. It appears to be impracticable to so preserve a true meat juice that it can become an article of commerce.

Miscellaneous Meat Preparations.—There is on the market a wide variety of manufactured products intended to replace beef juice. Some of these have meat extract as a base, and some have an addition of a small amount of albumin, or some form of soluble protein. Others consist largely of albumoses and peptones, and are formed by the action of steam or of acid and pepsin on meat. The tables on pages 247 and 248 give the composition of a number of products of this nature.

The preparations given in the table on page 248 are arranged in four classes, according to their content of proteoses and peptones, meat bases, creatin, and insoluble proteins.

Yeast Extract.—During recent years a product closely resembling meat extract has been prepared by the evaporation of the water extract of yeast. This product has been sold as a substitute for meat extract and has been reported in Germany as an adulterant therefor. The best means of distinguishing yeast extract from meat extract is by the determination of creatin and creatinin, which are absent in the former.*

Wintgen† has pointed out that the filtrate from the zinc sulphate precipitate obtained in the determination of albumoses is clear in the case of meat extracts, but turbid if a considerable percentage of yeast extract be present.

METHODS OF ANALYSIS.

Water.—Water is best estimated by weighing from 2 to 3 grams of the preparation (if of the dry or pasty variety), or from 5 to 10 grams of the fluid extract, into a large platinum dish, the dry variety being dissolved in a little hot water. The powdered preparations are dried directly without admixture. To pasty and fluid preparations are added sufficient ignited asbestos, pumice stone or sand, sifted free from dust, to absorb the solution. Pasty preparations are first dissolved in sufficient water to make them distinctly fluid. The sample is then dried at 100° C. till it ceases to lose weight. Tin or lead dishes or Hoffmeister glass dishes may be employed, and after being cut or broken, placed in the extraction tube for the determination of fat.

^{*} Micko, Zeits. Unters. Nahr. Genuss., 5, 1902, p. 193; 6, 1903, p. 781.

[†] Arch. Pharm., 242, 1904, p. 537.

· Partly insoluble.

MISCELLANEOUS PREPARATIONS (MEAT EXTRACTS, JUICES, AND POWDERS).*

	.beaimmstebaU	9 cm 44 km 0 = 0 000 km 0 000 m 000 0 m = 000 0 000 m 000 0 m = 000 0 000 m 000 0 m = 000 0 000 m 000 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	
	Ether Extract.	50000000000000000000000000000000000000	
	sinomnA	22222222222222222222	
	Meet bases Other than Creatin, Creatinin, and Xanthin.	4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4	
	Xanthin Bases.	000000000000000000000000000000000000000	
1	Creatin and Creatinin.	0 45 0 58 0 0 58 0 0 0 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	
Nitrogen	Total Meat Bases.	4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4	. 36.
Nit	Peptones.	HO4004H4400H04000	114. p.
	Proteoses,	400000 = F 7 40 F 9 10 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	Bal.
	Insoluble and Coagu- lable Protein.	20000000000000000000000000000000000000	Bur. of Chem., Bul. 114.
	Total Mitrogen.	NN - 00 WO NG H - 40 4 W 4 4 00 00 0 0 0 0 0 0 0 0 0 0 0	dur. of
Acidity.	As Lactic Acid	8 6 0 0 0 4 4 0 0 0 4 0 0 0 0 0 0 0 0 0 0	Agric, I
Aci	N/10 Sodium Hydroxide,, cc. per gram	0 400 0 40 0 000 = + = 4 4 8 8 9 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	8
ső.	orrespond Phosphoric Linesphoric		Dept
utuent	Organic Phosphoric	0000000100000 000 1 1 1 1 1 1 0 0 0 0 0	U S
Consi	Total Phosphoric Acid	4 4 4 0 4 0 0 4 0 0 0 0 0 0 0 0 0 0 0 0	Cook,
Mineral Constituent	Chlorine as Sodium Chloride in Ash,	2000000444 00 C 1 0 0 C 1 0 0 0 0 0 0 0 0 0 0 0 0	Bigelow and Cook,
	deA latoT	20 + + 0 4 + 0 4 + 4 0 + 8 8 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4	Bigelov
	.aetaW	4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4	•
		Royal Bouillon Capsules Bovril seasoned Beef jelly, Mosquera ext of beef beef beef beef Johnson's fluid beef (Bovril) tron American malted meat beef extract Arlington beef peptinoids	

Bigelow and Cook, U. S. Dept. of Agric, Bur. of Chem., Bul. 114, p. 26.
 b Largely coagulable, c Largely insoluble, d All coagulable.

a Best, mutton and fruits.

NITROGENOUS CONSTITUENTS OF MISCELLANEOUS PREPARATIONS.*

				1 0101	
g	AinominA	9202	22825481	48	F- 60
Nitrogen		6499	*******	5.0	>
i ii	_	445	44 C C C C C C C C C C C C C C C C C C	5.8	5855
Total	u K	2 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8	**********	4 0 4	
€		0.000	うちょうちょうちょう	4.4	8285
Terms of	Xanthin Bases.	0000	****		m000
, <u>5</u>		55.40	935 935 935 936	24	3 56 3 56 trace trace
	Creating and Creatinin	F-0-100	44-2 500	44	55
3		4400	80 AD 00 A00 00 A0 80 AD 07 PA D = A A0	F- 4	5467
Expressed in	Total Meat Bases.	3344	8 × 400 0 m = 4	4.5	2550
H		8250	=0.000.446	- 00 - 00	9.5
	Peptones.	9 1 1 4	200 00 4 0 0 0 000 00 4 0 0 0 000 00 00 00 00	55.0	204.
Bodies		<u> </u>			- POO 4
	Porteoses.	440 D 00	1 9 9 1 1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	4.00	as 46 ac =
8		4000	- wwg u r 4	€ →	9 7 7 7
1	TERROLAL MANUE	3483	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	W 101	4 4 9 4 4 60 0 0
Nitrogenous	-gao'd bas elduloen! saistore Proteins.	N = 4 4	400% 240	0 m	4886
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	летопи	0000	00000000	0.0	0000
'					
'	Creatin, Creatinin,	9229	30 00 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	300	8 9 7 6
'	Meat Bases Other than,	Natio	0 4 4 4 4 50 - 0	ın ó	~
		435	6-0 0 4-0 0 = 0	1 6	400- 500-
	Xanthin Bases.	0000	0000000	0.0	0000
_ ;		40.00	33. 53. 53.	9.9	5885
Bodies	ninitastO bas nitastO	H = = 0	6.00 1.31 0.10 0.61 0.63	0.0	o 47 o 25 trace trace
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5	Total Meat Bases.	00 0 m	0.0 P. 00 P. 0.0 P. 0.0	42 Or	E = □ □
Nitrogenous	,	1 20 0 2	000 = 4-0 × 4 4 400 0 4 0-00 = 4	900	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
🐉	Peptones.	P.48-		- 5	4 to 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
#			04055040	=	
-		2000	9995599	94	50.00
	Proteoses.	2022	04400000	0 M	4444
		<u> </u>	0 2 0 0 2 0 3	3 th	200 0 0 0 0 0 0 0 0 0
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	-aco3 has aldulosal	(1110	0 110 11 110 11	0 =	5 4 4 5
		9000	90760400	5.5	3 00 00 00
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l		328	7 2 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5	200	A P E A
		\ \text{A} \times \frac{1}{2} \t	ck incoming		Con Re
1		CLASS I Royal bouillon capsules. Boyril, seasoned Beef jelly Mosquera beef ext Brand & Co's essence of beef	Mulford's predigested beef. Armour's soluble beef Bovox essence of beef Johnson's fluid beef (Bovril) American brand beef ext Bovinine concentrated beef London Essence Co's ess of mutton Murdock's liquid food	Magg's bouillon Rose's peptonized best	CLASS IV Armour's beef ext and veg tablets. Loube-Rosenthal's beef solution American malted meat beef extract Arlangton beef peptinoids
1		ARRE	¥585583¥	ä	4744

† The sum of insoluble and coagulable proteins, proteoses, and peptones. Bigelow, and Cook, U. S. Dept. of Agric., Bur. of Chem., Bul. 114, p. 27

Ash.—From 2 to 3 grams of a fluid preparation, or a correspondingly less amount of a pasty preparation, are evaporated to dryness in a flat-bottom dish. Pasty preparations should first be dissolved in water, in order that the sample may distribute itself evenly over the bottom of the dish. The substance is then charred at the lowest possible heat, the charred mass exhausted with water, the insoluble residue collected on a filter and washed. The filtrate and residue are then returned to the dish and completely incinerated, the soluble portion of the ash added, evaporated to dryness, heated to a low redness and weighed. Chlorine is determined volumetrically or gravimetrically in the solution of the ash.

Fat.—This is best obtained by extracting a portion of the air-dried substance with petroleum ether in a Soxhlet apparatus. Petroleum ether extracts the fat only, while ether extracts other substances as well.

The determination is usually made in the residue from the determination of water. A properly prepared extract has very little fat.

Total Nitrogen.—The extract should be tested for nitrates, and the proper modification of the Gunning method should be employed, depending on the presence or absence of nitrates. Use from 1 to 5 grams for the determination. Nitrates should be properly accounted for.

Separation of Nitrogenous Compounds.—To correctly gauge the food value of a meat extract, it is essential to separate and estimate at least roughly its principal nitrogenous components. To attempt to make such a separation with a high degree of accuracy would involve a long and tedious series of operations, which in most cases would be impracticable. Usually the separation into three main groups is sufficient, insoluble proteins, tannin-salt precipitate (proteoses, peptones and gelatin) and meat bases. At times, however, it may become necessary, or at least desirable for specific purposes, to determine certain of the nitrogenous compounds separately.

Various quick methods have often been employed in connection with technical operations to determine the approximate amount of the several nitrogenous bodies or groups, but they have been generally discarded as untrustworthy. Among these may be mentioned Bruylant's* method of fractional precipitation by varying strengths of alcohol, and Hehner's† method of precipitation by methylated spirits. Another method that was widely used for a time was that of Allen and Searle,‡ which is based

^{*} Jour. Pharm. et Chem., 5, 1897, p. 515.

[†] Analyst, 10, p. 221.

[‡] Analyst, 22, 1897, p. 259.

on the belief that proteoses and peptones were completely precipitated from aqueous solution by saturating with bromine, after acidifying with hydrochloric acid. The experimental evidence on which this method was based consisted of the precipitation of proteins from the filtrate from the zinc sulphate precipitate, diluted with an equal volume of water. From the results so obtained, it appeared that peptones and proteins of larger molecule were completely precipitated by bromine in a half saturated solution of zinc sulphate, and it was assumed that precipitation from aqueous solution would be equally complete. Owing to a lack of methods by which peptones could be completely precipitated, this method has been widely used. The use of the method now appears to have been largely discontinued, as it has been repeatedly found to be unreliable.*

Complete Separation of Nitrogen Compounds would involve a discrimination between meat fiber and insoluble protein, coagulable proteins, acid albumin (syntonin), albumoses, peptones, meat bases, gelatin and ammonia.

- (1) Insoluble Proteins.—About 5 grams of the extract of the dry, or 20 to 25 grams of the fluid variety are exhausted with 200 to 250 cc. water at about 20° C., and the residue collected on a tared filter. It is often difficult to filter such an extract in the ordinary way, and the use of the centrifuge is helpful, passing the clear supernatant liquid through the filter, and finally washing the residue thereon. The residue is washed, dried at 100°, and weighed, or the nitrogen may be determined by the Gunning method. The sample may also be placed in a graduated flask, digested in a considerable amount of cold water for several hours with frequent shaking, and the nitrogen determined in an aliquot part of the filtrate. This deducted from total nitrogen gives the nitrogen of insoluble proteins. $N \times 6.25 = total$ insoluble matter, which includes, besides the meat fiber, the insoluble proteins.
- (2) Coagulable Proteins.—The filtrate from (1) is neutralized exactly to litmus, and dilute acetic acid added till acidity is just apparent. It is then boiled for some minutes to make insoluble the coagulable proteins, which are collected upon a filter (using to advantage a centrifuge as in the preceding paragraph). Determine the nitrogen in the washed residue, using the factor 6.25 for coagulable protein.
 - (3) Albumoses or Proteoses.†—An aliquot part of the filtrate from (2)

^{*} Bigelow, U. S. Dept. of Agric., Div. of Chem., Bul. 13, pt. 10, p. 1396; Bul. 81, p. 106. Sjerning, Zeits. anal. Chem., 39, 1900, p. 545. Fraps and Bizzell, Jour. Am. Chem. Soc., 22, 1900, p. 709. Van Slyke, Chem. News, 88, 1903, p. 92.

[†] Bömer, Zeit. anal. Chem. 5, 1895, p. 562.

is saturated with zinc sulphate, adding the powdered salt as long as it continues to dissolve with stirring and shaking. Proteoses and any traces of gelatin or insoluble proteins that have escaped removal are precipitated, but not the peptones or meat bases. Filter, wash, and determine the nitrogen in the residue, using the factor 6.25 for the proteoses.

(4) Peptones.—Sjerning's Tannin-salt Method, modified by Bigelow and Cook.*—An aliquot part of the filtrate from (2), concentrated by evaporation to 20 cc. or less, in case it is necessary to take more than 20 cc., is transferred to a 100-cc. flask.

Then 50 cc. of a solution containing 30 grams of sodium chloride per 100 cc. are added, and the flask agitated to insure the thorough mixing of its contents and the solution of the sample. The flask is now placed in the ice box at approximately 12° C. After the solution has reached the ice box temperature (this requires an hour usually), 30 cc. of a 24% solution of tannin (which must be at ice box temperature) are added. The total volume is now 100 cc. The contents of the flask are thoroughly mixed, and the flask returned to the ice box, where it remains over night. In the morning the solution is filtered at ice box temperature into a 50 cc. graduated flask. The nitrogen is determined in this filtrate, and also in an aliquot portion of the filtrate from a blank, in which the reagents alone are employed. The nitrogen found in the 50 cc. portion, multiplied by two (after correction for the nitrogen in the blank), gives the total nitrogen in the filtrate, and is calculated to per cent of nitrogen on the sample employed. This includes the nitrogen present as ammonia, and all of the nitrogen of the meat bases, except that portion of the creatin precipitated by the tannin-salt reagent. The figure thus obtained is added to the per cent of nitrogen as determined in (1), (2), and (3). This sum, deducted from the total nitrogen, is ordinarily reported as the per cent of nitrogen existing as peptones, and is multiplied by 6.25 for the per cent of peptones.

It is probable that the substances so reported are not true peptones, since the filtrate from (3) commonly gives no biuret reaction. They probably consist largely of peptoids, formed by the action of the hot solution on gelatin and polypeptides.

Bigelow and Cook find that the tannin-salt precipitate is not contaminated with other meat bases than creatin. They believe that about one-quarter of the creatin is found in this precipitate. Accordingly,

^{*} Jour. Am. Chem. Soc., 28, 1906, p. 1496.

they suggest that the percentage of creatin be determined before and after precipitation with tannin-salt reagent, and correction made by the results so obtained.

Street believes this correction to be impracticable. He finds that it is very difficult, if not impossible, to remove tannin completely from the filtrate, and that the slightest trace of tannin prevents the color reaction for creatin.

- (5) Meat bases.—The per cent of nitrogen found in the filtrate from the tannin-salt precipitate in (4), after deducting the per cent of nitrogen found as ammonia in (6), is multiplied by 3.12 to obtain the per cent of meat bases.
- (6) Ammonia.—From 5 to 10 grams of the original sample are dissolved in a convenient volume of water, and distilled after the addition of powdered magnesia. The distillate is titrated, and its alkalinity reported as per cent of NH₃. The corresponding percentage of nitrogen is also calculated, as it is necessary for the calculation of meat bases in (5).

Determination of Creatin and Creatinin.*—This determination may be made in an aliquot of the filtrate from the insoluble and coagulable protein determination.† The aliquot must contain sufficient total creatinin. after dehydration of the creatin to creatinin, to give a reading not far from 8° on the scale of the Dubosc colorimeter, after applying the colorimetric method as outlined by Folint for the estimation of creatinin in the urine. Heat this aliquot with 5 cc. of half-normal hydrochloric acid for three and a half hours on a steam bath under a reflux condenser. Neutralize the hydrochloric acid by the addition of 5 cc. of half-normal sodium hydroxide, then add 15 cc. of a saturated picric acid solution, and 5 cc. of 10% sodium hydroxid. Shake the solution, and allow it to stand for five minutes; make up to 500 cc., and compare the color with a half-normal solution of potassium bichromate in the Dubosc colorimeter. The half-normal bichromate solution when the scale is set at 8° corresponds to 10 mg. of creatinin, and from this figure the amount of creatinin in the aliquot is readily calculated.

Hehners criticises this method as applied to meat extracts. He believes that more complete results may be obtained by using 25 cc. of a 1.01% of picric acid with "a quite small amount of alkali." He considers the

^{*} Bigelow and Cook, Jour. Am. Chem. Soc., 28, 1906, p. 1497.

[†] Aliquot should represent approximately 0.2 gram of a first class solid beef extract.

[‡] Zeits. physiol. Chem., 41, 1904, p. 223.

[§] Pharm. Jour., 78, 1907, p. 683.

precipitate somewhat soluble in excess of alkali. Emmett and Grindley* have made an exhaustive study of the method as applied to meats, meat extracts, and urines. They find that 15 cc. of 1.2% picric acid should be employed for the original creatinin determinations, and 30 cc. for the dehydrated creatinin. They also recommend 5 cc. of alkali for the original creatinin, and 10 cc. for the dehydrated creatinin, though an additional 5 cc. does not give lower results.

Determination of Xanthin Bases.—In addition to creatin and creatinin, a true meat extract or meat juice should contain small amounts of xanthin bases, including xanthin, hypo-xanthin, guanin, and adenin. These bodies are derived from the nuclei of the cells, and, consequently, in an extract that is prepared from fresh, unaltered beef a certain amount of these bodies should be obtained, together with the salts and other extractive matter. The determination of the xanthin bases is, therefore, of value in determining the origin of an alleged extract of meat.

The xanthin base figures in the tables show a variety of results, which is explained by the fact that in the preparation of the extract under certain conditions of heat and pressure some of these bodies are destroyed. The following method was employed for their determination:

Schittenhelm's Method modified by Cook.†—Use an amount of the standard solution equivalent to 5 grams of the original extract. Place in a large evaporating dish, and add 500 cc. of 1% sulphuric acid. Evaporate to 100 cc. within 4 to 5 hours. Cool, and neutralize with sodium hydroxide. Add 10 cc. of 15% sodium bisulphate, and 15 cc. of 20% copper sulphate; allow this to stand over night, filter, and wash. The precipitate suspended in water is treated with sodium sulphide, and warmed on the steam bath. Add acetic acid to acidify, and filter hot. To the filtrate add 10 cc. of 10% hydrochloric acid, and evaporate to a volume of about 10 cc. Filter, make ammoniacal, and add ammoniacal silver nitrate of 3% strength. After standing several hours, the solution is filtered and the precipitate washed with distilled water until no longer alkaline. The nitrogen in the precipitate is that of the xanthin bases.

Determination of Gelatin.—This is accomplished by the modified Stutzer method as given on page 231.

Determination of Acidity.‡—In the average solid or pasty extract the lactic acid content varies from 4 to 8 per cent, and, as a rule, the extract showing the highest phosphoric acid content likewise shows the highest

^{*} Jour. Biol. Chem., 3, 1907, p. 491.

[†] U. S. Dept. of Agric., Bur. of Chem., Bul. 114, p. 41.

[‡] Ibid., p. 39.

acidity. This is undoubtedly due to the fact that some of the phosphoric acid is in the form of di-hydrogen or acid phosphate, although the character of the acidity has not been definitely determined.

The method employed for determining acidity consisted in adding tenth-normal sodium hydroxide to a dilute solution of the meat extract in water, until a drop removed by means of a small capillary tube and tested on a piece of litmus paper gives a neutral reaction. The results are expressed in cubic centimeters of tenth-normal sodium hydroxide, also as per cent of lactic acid present. The acidity is commonly expressed as per cent of lactic acid, though it is probably due in large part to acid potassium phosphates. Lactic acid is the chief organic acid, though succinic acid is also present in notable amount.*

Detection of Preservatives in Meat Extracts.—Boric acid is sometimes used as a preservative in these preparations, and is tested for by the usual methods (Chapter XVIII).

Determination of Glycerin.—This substance is sometimes used as a preservative for fluid preparations. Perhaps the most satisfactory method that has been suggested for its determination is that of Bigelow and Cook.† The dried residue is extracted with acetone, the meat bases removed by precipitation with silver nitrate, followed by phosphotungstic acid. The glycerin is determined in the filtrate by Hehner's method.†

FISH.

Structure and Composition.—Fish resembles meat both structurally and in the nature of its constituents, but differs from it in a marked degree in the relative proportions of its various components. Thus, there is considerably more refuse matter such as skin and bones in fish than in meat, and in the edible portion of fish the amount of water is much greater. Comparing the nitrogenous components of each, we find in fish more of the gelatin-yielding matter (collagen) and less of the extractives than in meat. There is much less hæmoglobin or allied coloring substance in the flesh and blood of fish than in meat, which accounts for the white color usually characteristic of the former. Certain fish, however, like the salmon, probably owe their distinctive color to a pigment belonging to the lipochrome § class. The mineral content of fish, as a rule, exceeds that of meat and contains more phosphates. The various edible fishes differ less among themselves in composition than do the meats. According to Chapman the average composition of fish is as follows, in parts per 1000:

^{*} Arb. kais. Gesundheitsamt, 1906, vol. 24.

[†] U. S. Dept. of Agric., Bur. of Chem., Bul. 114, p. 42.

[‡] Jour. Soc. Chem. Ind., 8, 1889, p. 4.

A series of fatty animal pigments.

Water	740.82
Albumin.	137.40
Collagen	
Fat	45-97
Extractives	16.97
Salts	14.06

Hutchison classifies fish as follows with reference to their content of fat: Lean.—Fish having less than 2% fat, such as cod and haddock.

Medium.—Fish having 2 to 5% fat, such as halibut and mackerel.

Fat.—Fish having more than 5% of fat, such as eel, 18%; salmon, 12%; turbot, 12%, and herring, 8%.

According to Atwater and Bryant * the composition of different varieties of fish is as follows:

COMPOSITION OF FISH.

				Pro	ein.			Fuel
		Refuse.	water.	N× 6.25.	By Di Ter- ence.	Fat.	Ash.	Value per Pound.
Bass-	edible portion		77-7	18.6	18.3	2.8	1.2	465
	as purchased.		35.1	8.4	8.3	I.I	0.5	200
Bluefish-			78.5	19.4	19.0	1.2	1.3	410
	as purchased	48.6	40.3	10.0	9.8	0.6	0.7	210
Cod—	edible portion	• • • • • •	82.6	16.5	15.8	0.4	1.2	325
T. 1	as purchased		38.7	8.4 18.6	8.0	0.2	0.6	165
Eel—	edible portionas purchased.		71.6		18.3	9.1	0.8	730
Waddaah	edible portion		57-2 81.7	14.8	14.6	7-2	1.2	580
Haddock-	as purchased.		40.0	8.4	8.2	0.3	0.6	335 165
Halibut	edible portion		75-4	18.6	18.4	5.2	1.0	565
manout—	as purchased.		61.9	15.3	15.1	4.4	0.9	470
Herring-	edible portion		72.5	19.5	18.g	7.1	1.5	660
	as purchased		41.7	11.2	10.0	3-9	0.0	375
Mackerel-	edible portion		73-4	18.7	18.3	7.1	1.2	645
	as purchased		40.4	10.2	10.0	4.2	0.7	365
Perch-	edible portion		75-7	19.3	10.1	4.0	1.2	530
	as purchased		28.4	7.3	7.2	1.5	0.4	200
Pickerel-	edible portion		79.8	18.7	18.6	0.5	1.i	370
	as purchased	47.I	42.2	9.9	10.7	0.3	0.6	210
Salmon-	edible portion		64.6	22.0	21.2	12.8	1.4	950
	as purchased		40.9	15.3	14.4	8.9	0.9	660
Shad—	edible portion		70.6	18.8	18.6	9-5	1.3	750
	as purchased	50.1	35.2	9.4	9.2	4.8	0.7	380
Skate—	edible portion		82.2	18.2	15.3	1.6	1.1	400
	as purchased		40.2	8.9	7.5	0.7	0.6	195
Smelt—	edible portion		79.2	17.6	17-3	1.8	1.7	405
_	as purchased		46.1	10.1	10.0	1.0	1.0	230
Trout—	edible portion		77.8	19.2	18.9	2.1	1.2	445
m	as purchased		40.4	9.9	9.8	1.1	0.6	230
Turbot—	edible portion		71.4	14.8	12.9	14-4	1.3	885
Whites	as purchased	47-7	37.3	7-7	6.8	7.5	0.7	460
w nitensn-	as purchased		69.8	22.9	22.1	6.5	1.0	700
	as purchaseu	53-5	32-5	10.0	10.3	3.0	0.7	325

^{*} U. S. Dept. of Agric., Off. of Exp. Sta., Bul. 28, p. 47 et seq.

Crustaceans and Mollusks:—These differ from the meats and common fish by reason of the presence in considerable proportion of the carbohydrate glycogen, contained in the liver. The *lobster* and *crab* are nearly alike in composition, the flesh being made up of coarse, dense, thick-walled fibers.

Payen gives the following composition of the flesh and body of lobster:

	Flesh (contained in Claws and Tail).	Body (consisting mainly of Liver).
Water	76.6	84.31
Protein	19.17	12.14
Fat	1.17	1.14

Clams and Oysters are low in solid nutriment, and are more digestible when eaten raw than cooked. Oysters contain 3% or more of glycogen.

The following analyses are from Atwater and Bryant:*

COMPOSITION OF SHELL FISH, ETC.

	Refuse.	Water.	Pro- tein. N× 6.25.	Fat.	Car- bohy- drates.	Ash.	Fuel Value per Pound. Cals.
Clams— edible portion		85.8	8.6	1.0	2.0	2.6	240
as purchased			5.0	0.6	1.1	1.5	140
Crabs— edible portion		77.I	16.6	2.0	1.2	3.1	415
as purchased	52.4	36.7	7.9	0.9	0.6	1.5	195
Lobster— edible portion		79.2	16.4	1.8	0.4	2.2	
as purchased	61.7	30.7	5.9	0.7	0.2	0.8	140
Mussels— edible portion		84.2	8.7	1.1	4.1	1.9	285
as purchased	46.7	44.9	4.6	0.6	2.2	1.0	150
Oysters— edible portion			6.2	1.2	3-7	2.0	235
as purchased		16.1	1.2	0.2	0.7	0.4	45
Scallops— as purchased			14.8	0.1	3-4	1.4	345
Terrapin—edible portion		74-5	21.2	3-5		1.0	545
as purchased	75-4	18.3	5.2	0.9		0.2	135
Turtle— edible portion		79.8	19.8	0.5		1.2	390
as purchased	76.0	19.2	4-7	0.1		0.3	90

Characteristics of Fresh Fish.—Fish of all kinds should be eaten when perfectly fresh, as it undergoes decomposition much sooner than meat when killed. While with meat aging is often beneficial to bring out requisite tenderness and flavor, in the case of fish deterioration begins almost immediately after death. Even though certain varieties of fish may be kept firm and wholesome for some days on ice, the flavor is distinctly impaired by long keeping. Fish that is not perfectly firm to the touch, or that has abnormally dry scales, or that shows blubber at the

^{*} U. S. Dept. of Agric., Off. of Exp. Sta., Bul. 28, pp. 52 and 53.

gills, or that possesses the marked odor that accompanies incipient decomposition, should not be used as food.

Methods of Analysis.—These are similar to the methods given for meat.

Preservatives in Fish and Oysters.—Boric acid and borax in mixture and sodium benzoate form the most common preservatives of salt dried fish and of oysters. In the case of salt codfish, the preservative is sprinkled on the surface. Such surface application in some states, as for example Massachusetts, is allowed by law. In opened oysters sold in casks and kegs, boric mixture has been used commonly in solution in the oyster liquor, but is now infrequent.

The author has found salicylic acid in bottled clam juice and clam bouillon.

CONCENTRATED FOODS.

Under the name of "condensed" or "concentrated foods" or "emergency rations" a number of canned preparations are sold for the use of campers, travelers, armies in the field, etc. These consist usually of mixtures of dried ground meats and vegetables, pressed together in compact form, and preserved in tin cans. The claims made for the food value of these preparations are, as a rule, extravagant and erroneous, as shown by Woods and Merrill,* who give the following analyses of some of these foods:

	Net Weight Con- tents.						
		Water.	Water. Pro-		Carbo- hy- drates.	Ash.	Fuel Value. Cals.
	Grams.	Grams.	Grams.	Grams.	Grams.	Grams.	
Ration cartridge, pea, beef, etc	241	.34.2	52.9	42.0	98.0	13.9	1071
Blue ration campaigning food, a	169	76.1	37-5	9.0	37.9	8.5	432
" " " b	78	10	5.6	23.1	46.9	1.4	436
Red ration campaigning food, a	122	33.8	26.2	18.5	37.8	5-7	496
" " b	77	1.2	5.0	23.0	46.6	1.2	424
Ration cartridge, potatoes, beef, etc	283	117.9	62.3	12.6	76.4	13.8	772
Emergency ration, a	120	14.2	56.1	29.6	11.9		617
« b	113	1.9	8.2	32.7	68.0	2.2	622
Emergency ration, a	121	4-5	71.8	32.6	6.7	5-4	776
" " b	127	5-7	8.3	15.3	94.8	2.9	588
Nao meat food	437	231.3	56.9	90.1	46.2	12.5	1328
Army rations	661	420.2	101.2	84.3	47-9	7.4	1542
Standard emergency ration	418	23.6	129.6	90.5	160.3	14.0	2198
" " a	270	17.0	50.6	54.8	137.0	10.6	1402
« « b	49	0.5	3.2	10.5	34.0	0.8	254
Arctic food		30.7	75.1	167.3	119.8	30.1	2430
Tanty emergency ration	475	313.5	60.2	48.6	41.9	10.8	1482
F-A Food Company's stew	964	638.0	149.2	114.5	52.5	9.8	2460

^{*} Maine Exp. Sta., Bul. 75, p. 103.

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CHAPTER IX.

EGGS.

Nature and Composition.—Though eggs of various birds are used to some extent as food, it is the egg of the hen that is in universal use for this purpose, and therefore the one which is here for the most part discussed, bearing in mind that the structure and composition of all varieties of birds' eggs are closely analogous.

Fig. 60 shows the longitudinal section of a hen's egg.

Fig. 60.—Longitudinal Section of a Hen's Egg. 6, Shell; b, Double Membrane of Shell; c, Air-chamber; d, Outer, or Fluid Albuminous Layer; e, Thick, Middle Albuminous Layer; f, Inner Albuminous Layer; g, Membrane of the Chalaza; hh, the Chalaza; i, Vitelline Membrane; j, Germ; k, Yolk; l, Latebra. (After Macé.)

The average weight of a hen's egg is 60 grams, of which the shell weighs about 6, the white 36, and the yolk 18. Roughly it contains 70% of water, 12% of albumin, and 12% of fat.

The shell, according to König, has the following composition:

Calcium carbonate	89-97%
Magnesium carbonate	0- 2%
Calcium and magnesium phosphate	0.5- 5%
Organic substances	2.0- 5%

The mean percentage composition of the eggs of the hen	, duck,	and
plover are, according to König, as follows:		

	Water.	Proteins.	Pat	Pat, free Sub- er Cent.	Salts. Per Cent.	In the Dry Sub- stance.		
	Per Cent.		Per Cent.			Nitrogen Per Cent.	Fat Per Cent.	
Hen's egg	73.67	12.55	12.11	0.55	1.12	7.66	45.99	
Duck's egg	71.11	12.24	15.49		1.16	6.78	53.62	
Plover's egg	74-43	10.75	11.66	2.18	0.98	6.75	45.78	
White of hen's egg	85.75	12.67	0.25		0.59	14.25	1.78	
Yolk " " "	50.79	16.24	31.75	0.13	1.00	5.30	64.43	

The Egg-white.—The white of egg has a specific gravity of 1.045, and its reaction is always alkaline. It is a transparent, albuminous fluid inclosed in a framework of thin membrane. The fibrous portion of the membrane is insoluble in water and in dilute acetic acid.

The composition of the fluid substance of the white of egg, according to Lehmann, is as follows:

Water	82 to 88	3%
Solids	13.3%	(mean)
Proteins	12.2%	46
Sugar	0.5%	44
Fats, alkaline soaps, lecithin, cholesterin	traces	
Inorganic residue	0.66%	,)

The protein substance is for the most part albumin, with a small amount of globulin.

According to Osborne and Campbell * the nitrogen compounds of the white of egg are four in number, which they name ovalbumin, ovomucin, conalbumin, and ovomucoid. No sharp and distinct separation of these bodies has yet been made.

Ovalbumin (albumin) is the chief constituent, and forms by far the largest portion of the protein of the egg-white. In 2.5% solution in water, ovalbumin starts to coagulate at 60°, and yields a dense coagulum at 64°. Stronger solutions require a somewhat higher temperature for coagulation.

Ovomucin is a globulin-like substance, precipitated from egg-white by dilution with water. It is partly soluble in strong sodium chloride solution. When dried and washed with alcohol, it is a light white powder.

Conalbumin bears a close resemblance to ovalbumin, but coagulates

^{*} Jour. Am. Chem Soc., 22 (1900), p. 422.

EGGS. 263

in dilute salt solution at a lower temperature (below 60°), and the coagulum is more flocculent than that of ovalbumin.

Ovomucoid is not coagulable by heat, and may thus be separated (imperfectly) by filtering out all the coagulable proteins.

The last two compounds exist in very small amounts only.

Preparation of Albumin.*—By beating up the white of egg in water, the salts and the albumin are dissolved, while the fibrous portion is insoluble and is removed by filtration. The filtrate is then treated with a slight excess of basic lead acetate, the precipitate decomposed by treatment with carbon dioxide, and the lead removed by hydrogen sulphide. The solution is warmed cautiously to 60° C., thus beginning to coagulate the albumin, a small part of which, coming down in a flaky form, carries with it the lead sulphide. On filtering or pouring off the supernatant liquid after cooling, one obtains a colorless solution of the albumin, which is evaporated to dryness below 40°. The albumin is obtained in the form of transparent yellowish, horny scales, which may be pulverized in a mortar, if desired. Its specific gravity is 1.262. It is tasteless, odorless, and neutral in reaction, and slowly soluble in water.

The Egg-yolk.—This is much more complex in composition than the white. Halliburton thus enumerates the constituents of the yolk:

(a) Proteins.—Vitellin, the chief one, a globulin resembling myosin. Albumin, in small quantities.

Nuclein, combined chiefly with the iron present.

(b) Fats.—Olein, palmitin, and stearin.

A yellow lipochrome or lutein.

- (c) Carbohydrates.—Grape sugar in small quantities.
- (d) Other Organic Constituents. Lecithin, a phosphorized nitrogenous body allied both to the fats and to the proteins.

Cerebrin.

Cholesterin.

(e) Inorganic Salts, the most abundant of which is potassium chloride. Gobley gives the following composition to the egg-yolk:

F	Per Cent.	Per	Cent.
Vitellin	1.5 0.3 7.2	Cholesterin. Fats. Coloring matters. Salts. Water.	20.3 0.5 1.0

^{*} Allen, Com. Org. Anal., Vol. IV, p. 42.

Osborne and Campbell,* as the result of long and careful experiments, consider the protein of egg-yolk to be largely if not wholly a lecithin compound, having properties of a globulin, and soluble in sodium chloride solution.

The fat of the egg yolk, which is used in ointments, has the following characteristics according to Spaeth: †

Specific gravity at 100° C	0.881
Iodine number	68.48
Reichert-Meissl value	0.66
Refractive index at 25° C. (on butyro-refractometer scale)	68.5
Melting-points of fatty acids	36° C.
Iodine number of fatty acids	72.6

The mineral content of the egg is thus shown by König:

COMPOSITION OF THE ASH OF EGGS.

		Ash of the Dry Sub- stance.	Potash.	Soda.	Lime.	Mag- nesia.	Iron Oxide.	Phos- phoric Acid.	Sul- phuric Acid.	Silica.	Chlo- rine.
Hen's egg:	white	4.61	17.37 31.41 9.29	31.57	2.78	2.79	0.57	4.41	2.12	1.06	8.98 28.82 1.95

The following analyses of eggs were made by Wood and Merrill: ‡ AVERAGE WEIGHTS OF EGGS AND PARTS AS PREPARED FOR ANALYSIS.

	Weight		Weight	Boiled.	Shell	Whi.			
	as Received.	Shell (Refuse).	White.	Yolk.	Total.1	(Refuse).	White.	Yolk.	
	Grams.	Grams.	Grams.	Grams.	Grams.		Per Cent.	Per Cent.	
Turkey	105.5	24.1	60.1 98.5	30.9 64.8	102.7	11.4	56.5 52.6	30.1	
Goose Duck	70 6	7.2	36.5	24.4	68.1	10.6	53.6	34.0 35.8	
Guinea fowl	40.2	5.6	20.9	12.5	39.0	14.4	53.6	32.0	

¹Shrinkage due to loss in preparation and cooking.

^{*} Jour. Am. Chem. Soc., XXII, 1900, p. 413.

[†] Abst. Analyst, 1896, p. 233.

[‡] Maine Exp. Sta., Bul. 75, p. 90.

COMPOSITION OF EGGS.

				Prot	ein.			ig.
		Refuse (Shells).	Water.	Nitrogen ×6.25.	By Dif. ference.	Fat.	Ash.	Fuel Value per Pound.
Turkey—	whiteyolk. entire edible portionas purchased.	13.8	86.7 48.3 73.3 63.5	11.5 17.4 13.4 11.6	12.5 17.6 14.2	Trace 32.9 11.2 9.7	o.8 1.2 0.9	Cal. 325 1875 850
Goose-	white	14.2	86.3 44.1 69.5	11.6 17.3 13.8	12.9 18.4 15.1	Trace 36.2 14.4 12.3	0.8 1.3 1.0	735 330 1975 985 860
Duck-	whiteyolkentire edible portionas purchased.		87.0 45.8 70.5 60.0	11.1 16.8 13.3	12.2 16.8 14.0	Trace 36.2 14.5	0.8 1.2 1.0 0.8	315 1980 985 880
Guinea fowl-	whiteyolkentire edible portionas purchased	16.9	86.6 49.7 72.8 60.6	11.6 16.7 13.5 11.2	12.6 17.3 14.3 11.0	Trace 31.8 12.0	0.8 1.2 0.9	325 1800 875 730
Hen—	whiteyolkentire edible portionas purchased	11.2	86.2 49.5 73.7 65.5	12.3 15.7 13.4 11.9	13.0 16.1 14.8 13.1	0.2 33.3 10.5 9.3	0.6 1.1 1.0 0.9	

METHODS OF ANALYSIS.

Preparation of the Sample.*—The egg is first weighed as a whole and afterwards boiled hard, cooled, and again weighed. The shell, white, and yolk are then carefully separated and each weighed. After rejecting the shell, the yolk and white are separately reduced by a chopping-knife to the size of wheat grains. These portions are dried partially at a temperature not exceeding 45°, weighed, and afterwards ground to a fine powder in a mortar.

Determinations of water, fat, ash, and total nitrogen are made in practically the same manner as with flesh foods.

Little attention has been paid as yet to the complete separation and determination of the nitrogen compounds in the white and yolk, and it is customary in most cases to express the protein of the whole as N×6.25.

Determination of Lecithin.—Wiley's Method.†—The whole egg, excluding the shell, is placed in a flask with a reflux condenser, and boiled for six hours with absolute alcohol. The alcohol is then evaporated off, and the residue treated in like manner for ten hours with ether. After evaporat-

^{*} Woods and Merrill, Maine Exp. Sta., Bul. 75, p. 92.

[†] Principles and Practice of Agricultural Analysis, Vol. III, p. 431.

ing off the ether, the dry residue is rubbed to a fine powder, placed in an extractor and treated with pure ether for ten hours. The ether extract thus secured is oxidized, after removal of the ether, by fusion with mixed sodium and potassium carbonates, and the phosphorus is determined in the usual way as magnesium pyrophosphate. The amount of lecithin is obtained by multiplying the weight of magnesium pyrophosphate by the factor 7.2703, on the basis of Hoppe-Seyler's formula for lecithin: $C_{44}H_{80}NPO_{8}$.

If, for example, an amount of organic phosphorus yielding 0.0848 gram of magnesium pyrophosphate is found in 54 grams of egg exclusive of shell, then $0.0848 \times 7.2703 = 0.61652$ and $0.61652 \times 100 \div 54 = 1.14$. Therefore the percentage of lecithin in the egg is 1.14.

Preservation of Eggs.—Owing to the porous nature of the shell, the moisture of the contents gradually grows less by evaporation, and the egg loses in weight. Air also passes in through the shell pores, carrying various microbes, which result in ultimate decomposition and spoiling of the egg. Nature has provided the shell with a thin surface coating of mucilaginous matter, which, however, is easily washed off. This coating tends to partially close the pores, and for best results in keeping should not be removed by washing.

Eggs are commonly preserved by protecting them as far as possible from the air. This is accomplished in a variety of ways, the most common being to pack the eggs in salt or bran, so that the packing medium fills up the interstices between the eggs. Eggs thus packed will keep considerably longer then when exposed to the air. A solution of salt is sometimes employed, and also lime water, the eggs being simply packed in the solution. The use of lime water is, however, open to the serious objection that a disagreeable odor and taste are imparted to the eggs.

Eggs are sometimes coated with gelatin, vaseline, wax, or gum, so as to cover them with an impervious layer, either by dipping them in the coating medium, or by varnishing or otherwise applying the substance to the egg shell. By far the most efficacious egg coating has been shown by experiments in the North Dakota Experiment Station,* and also in Germany, to be sodium and potassium silicate, or water glass. The fresh eggs, preferably unwashed, are packed in a jar, and a 10% solution of water glass is poured over them. According to the North Dakota experiments, at the end of three and a half months, eggs packed in this manner the first of August appeared to be perfectly fresh.

^{*} Farmer's Bul. 103, U. S. Dept. of Agric., p. 18.

EGGS. 267

One drawback to this method is that eggs so treated break more easily on boiling, but this may be prevented by carefully piercing the shell with a strong needle.

Cadet de Vanx has proposed immersing the egg in boiling water for twenty seconds, the result being that a very thin layer of the egg-white next the shell becomes coagulated, thus forming an impervious coating inside the shell.

Cold-storage Eggs.—The preservation of eggs by storage at low temperatures has become an enormous industry. The temperature employed varies from 24° to 40° C., and the length of storage from one to eight months.

Experiments conducted by Wiley,* under authorization from Congress, have brought out certain points as to the physical and chemical changes that take place during cold storage. After breaking the shell and keeping at room temperature one day, the odor of eggs stored for 3.5 months was different from that of fresh eggs, but was not disagreeble. This odor increased on longer storage, and after 12.6 months became very characteristic. After 16.6 months, a musty odor was noticed immediately after opening the egg.

Chemical analysis by Cook showed that eggs in storage for one year lost 10% of the total weight, due to evaporation of water from the whites. Storage also caused a lowering of the amount of coagulable protein and of lecithin phosphorus, but an increase in lower nitrogen bodies, proteoses, and peptones. The acid reaction of yolks diminished during storage.

Microscopical examination by Howard and Read brought out the interesting fact that small rosette crystals of an unidentified substance appeared in the yolk after storage for 12 months or longer, and this observation has since been utilized in the examination of suspected samples.

Physical Examination of Eggs.—Various physical tests have been prescribed for ascertaining the approximate age of an egg. Thus, according to Delarne, if the egg, when placed in a 10% salt solution, sinks to the bottom, it may be considered perfectly fresh; if it remains immersed in the liquid, it is to be considered at least three days old; and if it rises to the surface and floats thereon it is more than five days old. This test

^{*} U. S. Dept. of Agric., Bureau of Chem., Bul. 115.

is a very rough one, and is useful only for eggs that have been kept in the air. Preserved eggs cannot be gauged by this means.

The best method of examining eggs for freshness is "candling," consisting in placing the egg between a bright light and the eye. If the egg is fresh, it will show a uniform rose-colored tint, without dark spots, the air-chamber being small and occupying about one-twentieth the capacity of the egg. If the egg is not fresh, it will appear more or less cloudy, being darker as the egg grows older, becoming in extreme cases opaque. At the same time the air-chamber grows larger as the age increases. So-called "spots" are eggs which show on candling black patches due to fungi.

Opened Eggs.—In the handling of eggs many become cracked or otherwise injured to an extent which renders them unfit for transportation. These are either sold to bakers for immediate use, or else opened and kept from spoiling by freezing, the addition of preservatives, or drying. The portions of "spot eggs" that do not show evidence of damage are also treated by one of these methods. Eggs which, because of their offensive taste, are unfit for food, are used in the tanning industry.

Preservatives commonly employed in opened eggs are boric acid and formaldehyde. The latter is especially effective as an egg preservative. If a small quantity be added and stirred into opened eggs that have become absolutely putrid, the result is astonishing. The product is completely deodorized, and exhibits the outward appearance at least of fresh eggs.

Formaldehyde, if present, may readily be detected by heating some of the egg directly with the hydrochloric-acid ferric-chloride reagent used in testing milk for formaldehyde, carrying out the process exactly as in the case of milk.

Desiccated Egg.—It is possible to evaporate to dryness the contents of the egg to form a powder, the keeping qualities of which far exceed that of ordinary eggs, while it forms a concentrated food which lends itself much more readily to transportation than does the fresh egg in the shell. Several brands of desiccated egg are on the market, which from their analyses are undoubtedly genuine. The following are analyses of two of them, one (A) made by the Bureau of Chemistry, the other (B) by the Massachusetts State Board of Health:

	A.	В.
Water	6.80	5-95
Protein (N×6.25)	45.20	48.15
Protein by difference	51.20	
Fat	38.5	40.56
Ash	3.5	5.34

Egg Substitutes.—There have been many preparations in powdered form sold under this name, nearly all claiming to contain all the ingredients of eggs, but most of them falling far short of these claims. Some of them, as for instance those made from desiccated skimmed milk, do contain nitrogenous matter, but as a rule little if any fat.

Two samples of "egg substitute" sold in Massachusetts were analyzed with the following results:*

	A.	В.
Protein	. 16.94	18.72
Fat		3.40
Water	. 6.71	7.01
Corn-starch, salts, and color	r-	
ing matter	- 72.92	70.87

A ten-cent package of sample A, weighing about 2 ounces, was alleged to be equivalent to 12 eggs. Starch furnished the chief ingredient in both samples.

One of the most flagrant examples of fraud in this connection was a product sold under the name "N'egg," advertised to contain the nutritive equivalent of the whites and yolks of a dozen eggs, "their composition being based on careful scientific analysis of natural eggs." It was put up in two small boxes, one containing a white and the other a yellow dry powder. Both were entirely devoid of nitrogen, and consisted of nearly pure tapioca starch with a little common salt, the color of the "yolk" being due to Victoria yellow.

Some egg substitutes are sold under the name of "custard powders," and are alleged to take the place of eggs in cooking. These are variously made up of mixtures of skim-milk powder, coloring matter, and baking powder ingredients as shown from the following analyses:†

^{*} An. Rep. Mass. State Board of Health, 1895, p. 675.

[†] Food and Sanitation, Nov. 25, 1893.

CUSTARD POWDERS.

	τ	2	3	4	5.	6
Starch	86.25	84.45	51.03	26.38	52.32	53.82
Albuminous compounds	0.59	0.58	6.01	2.96	6.00	5.06
Soluble coloring matter	0.88	0.98		,		"
Baking soda			15.33	50.70	22.11	26.71
Tartaric acid			13.69	10.33	11.37	6.10
Phosphates			0.24			_
Carbonates of lime and magnesia	• • • • •		2.70	'		
Chlorides and sulphates						1
Water.	11.83	13.69	11.00	9.63	8.20	8.22
Ash.	0.45	0.38		ا آ	l	

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" 103. Preserving Eggs.

CHAPTER X.

CEREALS AND THEIR PRODUCTS, LEGUMES, VEGETABLES, AND FRUITS.

THE chief points of difference in composition between the animal foods already treated of, and those of the vegetable kingdom, are apparent in the relative amounts of proteins and carbohydrates. The proteins present in the cereals and vegetables differ materially both in character and amount from those in the flesh foods, being as a rule present to a much greater extent in the meats than in the grains and vegetables. The leguminous foods, such as peas, beans, and lentils, are, however somewhat exceptional in this respect, being comparatively high in nitrogenous content.

The carbohydrates, which in the flesh foods are almost entirely lacking, and in milk make up about one-third of the solid matter, form the most important and abundant class of constituents in the vegetable foods.

The composition of the principal cereal grains is tabulated as follows by Villier and Collin:

	Wheat.	Barley.	Rye.	Oats.	Rice.	Corn.	Millet.	Buck- wheat.
Water	13.65	13.77	15.06	12.37	13.11	13.12	11.66	12.93
Nitrogenous substances.	12.35	11.14	11.52	10.41	7.85	9.85	9.25	10.30
Fat	1.75	2.16	1.79	5.32	o.88	4.62	3.50	2.81
Sugar	1.45	1.56	0.95	1.01	ן	2.46	1)	
Gum and dextrin	2.38	1.70	4.86	1.79	16.52	3.38	65.95	55.81
Starch	64.08	61.67	62.00	54.08		62.57	[] · · · [
Cellulose	2.53	5.31	2.01	11.19	0.63	2.49	7.29	16.43
Ash	1.81	2.69	1.81	3.02	1.01	1.51	2.35	2.72

The following results of the analyses of cereal grains are summarized from the work of the Division of Chemistry, United States Department of Agriculture:*

CEREAL GRAINS.

	Num- ber of Analy- ses.	Weight of 100 Ker- nels, Grams.	Moist- ure.	Pro- teins.	Ether Ex- tract.	Crude Fiber.	Ash.	Carbohydrates, other then Crude Fiber.	Wet Gluten.	Dry Gluten.
Barley:	14									
Mean		4-533	6.47	11.52	2.67	3.81	2.87	72.66		
Buckwheat:	10	. 555			•					
Mean		3.069	12.31	10.86	2.06	10.57	1.85	63.34		
Corn, domestic:		"	-			"	- 1			
Maximum		48.312	12.32	11.55	5.06	2.00	1.55	75.07		
Minimum		10.608	9.58	8.58	2.94	1.00	1.19	68.97		
Mean		38.979	10.93	9.88	4.17	1.71	1.36	71.95		
Oats, domestic:		1		-	-					
Maximum		3.891	13.02	15.05	6.14	16.65	4-37	61.44		
Minimum		2.038			0.93	8.57	2.47	53.70		
Mean		2.918	10.06	12.15	4.33	12.07	3.46	58.75		
Rice:				_				·		
Unhulled	4	2.929	10.28	7-95	1.65	10.42	4.09	65.60		
Unpolished	6	2.466	11.88		1.96	0.93	1.15	76.05		
Polished	14	2.132	12.34	7.18	0.26	0.40	0.46	79.36		
Rye, domestic:										
Maximum		4.201	11.45	18.99	2.30	2.50		75.36		
Minimum			9.54			1.65	1.71	63.61		
Mean		2.493	10.62	12.43	1.65	2.09	1.92	71.37		
Wheat, domestic:		l								
Maximum		6.190	14-53	17.15	2.50	3-72	2.35	76.05		
Minimum		2.125			0.28	1.70	1.40	66.67		4.70
Mean	• • • • •	3.866	10.62	12.23	1.77	2.36	1.82	71.18	26.46	10.31
Wheat, foreign:		ł				_		_	1	ł
Maximum			12.97			2.89	2.04		32.57	
Minimum		2.250		8.58		1.87	1.67	67.01		
Mean		4.076	11.47	12.08	1.78	2.28	1.73	70.66	25.36	9.82

Balland * gives the following percentage composition of beans, lentils, and peas:

	Bes	ans.	Len	tils.	Peas.		
	Min.	Max.	Min.	Max.	Min.	Max.	
Water. Nicrogenous substances. Fat. Sugars and starches. Cellulose. Ash.	10.10 13.81 0.98 52.91 2.46 2.38	20.40 25.46 2.46 60.98 4.62 4.20	11.70 20.42 0.58 56.07 2.96	13.50 24.24 1.45 62.45 3.56 2.66	10.60 18.88 1.22 56.21 2.90 2.26	14.20 22.48 1.40 61.10 5.52 3.50	

^{*} Jour. Pharm. Chem., 1897, pp. 196, 197.

The composition of potatoes, according to Balland,* is as follows:

	Water.	Nitroge- nous Sub- stances.	Fat.	Sugar and Starch.	Cellulose.	Ash.
Normal state—minimum maximum Dried— minimum maximum	66.10 80.60	1.43 2.81 5.98 13.24	0.04 0.14 0.18 0.56	15.58 29.85 80.28 89.78	0.37 0.68 1.40 3.06	0.44 1.18 1.66 4.38

The composition of the common vegetables, fruits, and berries is thus given by Atwater and Bryant.†

VEGETABLES.

		Number of Analyses.	Refuse.	Water.	Protein.	Fat.	Total Carbo- hydrates.	Crude Fiber.	Ash.	Fuel Value per Pound, Calories.
Asparagus—	as purchased	3		94.0	1.8	0.2	3.3	.8	-7	105
Beans, dried—	as purchased	11		12.6	22.5	1.8	59.6	4.4	3.5	1605
Beans, fresh Lima-		ī		68.5	7.1	.7	22.0	1.7	1.7	570
Dealis, it can Lima	as purchased	_	55.0	30.8	3.2	3	9.9	1.8		255
Beets, fresh-	edible portion	24	1	87.5	1.6	.1	9.7		1.1	215
beets, fresh-		24				1		.9		
0.11	as purchased		20.0	70.0	1.3	1.	7.7		-9	170
Cabbage—	edible portion	16		91.5	1.6	•3	5.6	1.1	1.0	145
	as purchased		15.0	77-7	1.4	.2	4.8		-9	125
Carrot, fresh-	edible portion	18		88.2	1.1	•4	9-3	I.I	1.0	210
	as purchased		20.0	70.6	-9	.2	7.4		-9	160
Celery—	edible portion	5		94-5	I.I	-1	3.3		1.0	85
	as purchased		20.0	75.6	.9	.I	2.6		.8	70
Cauliflower—	as purchased	2		92.3	1.8	-5	4-7	1.0	-7	140
Cucumber—	edible portion	4		95.4	8.	.2	3.1		••5	8o
	as purchased		15.0	81.1	-7、	.2	2.6		-4	70
Lettuce-	edible portion	8	l	94.7	1.2	-3	2.9	-7	.9	90
	as purchased	l	15.0	80.5	1.0	.2	2.5		.8	75
Mushrooms-	as purchased	11		88.1	3-5	-4	6.8	.8	1.2	210
Onion, fresh-	edible portion	15		87.6	1.6	-3	9.9	.8	.6	225
0111011, 1110011	as purchased	-3	10.0	78.0	1.4	.3	8.9		-5	205
Parsnip	edible portion	3	10.0	83.0	1.6	-5	13.5	2.5	1.4	300
Luismp	as purchased		20.0	66.4	1.3	.4	10.8		1.1	240
Pumpkin-	edible portion	3	20.0	93.1	1.0	.1	5.2	1.2	.6	120
1 umpkiii—	as purchased		50.0	46.5		.1	2.6		-3	60
Radish—	edible portion		30.0	01.8	-5	l .	8.3		1.0	
Radisii—	as purchased	4		1 4	1.3	-3	1 %	-7		135
Rhubarb-			30.0	64.3	-9	.I		-7	•7	95
Knubaro—	edible portion	2		94.4	.6	-7	3.6	1.1	-7	105
	as purchased		40.0	56.6	-4	-4	2.2		-4	65
Squash—	edible portion	10		88.3	1.4	-5	9.0	.8	.8	215
m	as purchased	-	50.0	44.2	-7	.2	4.5		•4	105
Tomato, fresh-	as purchased	27		94.3	-9	-4	3.9	.6	·5 .8	105
Turnip—	edible portion	19		89.6	1.3	.2	8.1	1.3	.8	185
	as purchased		30.0	62.7	-9	.1	5-7		.6	125
			I		Į	Ī	ļ	1		1

^{*} Jour. Pharm. Chem., 1897, pp. 298-300.

[†] Bul. 28, Office of Exp. Station U. S. Dept. of Agriculture.

FRUITS.

			KOIL	<u>.</u>						
		Number of Analyses.	Refuse.	Water.	Protein.	Fat.	Total Carbo- hydrates.	Crude Fiber.	Ash.	Fuel Value per Pound, Calories.
Apples—	edible portion			84.6		-5	14.2		•3	290
Apricots—	edible portion	11	25.0	85.0		•3	13.4	••••	·3	270
Bananas—	as purchased	6	6.0	75.3	1.3	.6		1.6	.8	²⁵⁵ 460
Blackberries-	as purchased	9	35.0	86.3		1.0	14.3	2.5	.6 •5	270
Cherries—	edible portion	16	5.0	80.9 76.8	1.0	.8			.6 .6	365 345
Cranberries— Currants—	as purchased	3		88.9 85.0	-4 1.5		9.9 12.8		-2 -7	215 265
Figs, fresh— Grapes—	as purchased edible portion	28 5		79.1 77.4	1.5	1.6	18.8 19.2	4-3	.6	380 450
Huckleberries-	as purchased -edible portion		25.0		1.0	1.2	14.4		•4	335 345
Lemons—	edible portion	4	30.0	89.3		·7	8. ₅	1.1	•5	205
Muskmelons-	edible portion	1	50.0	89.5	ð.		9.3 4.6	2.I	.6	185
Oranges-	edible portion	23	27.0	86.9	.8	.2 .1	11.6 8.5		•5 •4	240 170
Pears—	edible portion	2	10.0	84.4	.6	•5 •4	14.1	2.7	•4	295
Pineapple— Plums—	edible portion	1 3		89.3 78.4	-4	•3	9-7		-3	200 395
Prunes—	as purchased	24	5.0		.9		19.1		•5	370
Raspberries—	as purchased		5.8	75.6	-7		17.4	2.0	•5	335
Strawberries—				85.8 90.4	1.0	.6	7.4	1.4	.6	255 180
Watermelon-	edible portion as purchased	2	5.0	92.4	-4	.6 .2		••••	.0 .3	175 140 60
	m harmana		59.4	37-5	.2		2-/	• • • • •	٠.	~

The following analyses of apples made by Browne* are of interest. The first four analyses show the changes that occur in the composition of a Baldwin apple at different stages of its growth. Below these is given the average of the analysis of 160 samples, representing 27 varieties of apples.

COMPOSITION OF A BALDWIN APPLE AT DIFFERENT PERIODS.

Condition.	Water.	Solids.	Invert Sugar.	Su- crose.	Total Sugar.	Total Sugar after In- version.	Starch.	Free Malic Acid.	Ash.	Sugar Co- efficient.
Very green Green Ripe Over-ripe	79.81 80.36	18.47 20.19 19.64 19.70	6.40 6.46 7.70 8.81	1.63 4.05 6.81 5.26	8.03 10.51 14.51 14.07	8.11 10.72 14.87 14-35	4.14 3.67 0.17	1.14 0.65 0.48	0.27 0.27 0.28	47.16 53.10 75.71 72.84

^{*} Penn. Dept. of Agriculture, Bulletin 58.

AVERAGE COMPOSITION OF 27 VARIETIES OF APPLES.

Water	83.57
Solids	16.43
Invert sugar	7-92
Sucrose	3-99
Total sugar	11.91
Total sugar after inversion	12.12
Free malic acid	0.61
Ash	0.27
Sugar coefficient	73.76

The composition of the commoner nuts is shown in the following table:*

NUTS.

		Number of Analyses.	Refuse.	Water.	Protein.	Fat.	Total Carbo- hydrates.	Crude Fiber.	Asb.	Fuel Value per Pound Cal's.
Almonds—	edible portion. as purchased	11	45.0	4.8	20.0	54·9 30.2	17-3 9-5	2.0	2.0 I.I	3030 1660
Beechnuts-	edible portion. as purchased	I	40.8	4.0	21.9	57·4 34.0	13.2		3·5 2·1	3075 1820
Brazil-nuts	edible portion. as purchased	1	49.6	5.3	17.0	66.8 33-7	7.0 3.5		3.9	3265 1655
Butternuts-	edible portion. as purchased	I	86.4	4-4	27.9 3.8	61.2	3 5		2.9	3165
Chestnuts, fresh-	edible portion.	9	16.0	45.0	6.2	5.4	42.1 35.4		1.3	1125
Cocoanuts-	edible portion. as purchased	1	48.8	14.1	5.2 5.7 2.0	50.6 25.9	27.9 14.3		1.7	945 2760
Filberts—	edible portion. as purchased	1	52.I	3.7	15.6	65.3	13.0		.9 2.4	3290
Hickory-nuts-	edible portion. as purchased	1	62.2	3.7	7·5 15·4 5.8	31.3 67.4	11.4		1.I 2.I	1575 3345
Peanuts-	edible portion.	4		9.2	25.8	25.5 38.6	4·3 24·4	2.5	2.0	1265 2560
Pecans-	as purchased edible portion.	1	24.5	3.0	19.5	29.1 71.2	18.5		1.5	1935 3455
Pistachios—	as purchased edible portion.	1	53-2	4.2	5.2	33·3 54.0	6.2 16.3		3.2	1620 2995
Walnuts, Calif'nia-	as purchased		73-1	2.5 .7	18.4	17.3	13.0 3.5	1.4	1.7 -5	3306 885

Vegetables and Fruits furnish a large and most important portion of our food supply, but are naturally not included in their fresh state among the foods examined by the public analyst for adulteration, hence

^{*} U. S. Dept. of Agric., Off. of Exp. Station, Bul. 28.

but little space need be given them beyond a résumé of their composition, and an outline of methods of proximate analysis applicable to their examination for food values. When, however, these products undergo the various processes incidental to their treatment for long keeping, such as preserving, canning, drying, pickling, and mixing with other ingredients, it is then that many varieties of fraudulent adulteration are practiced. Vegetable foods thus prepared form the subject of a separate chapter. Besides the proximate components that commonly occur in vegetable products, there are three other substances worthy of mention found in vegetables and fruits, viz., inosite, pectose, and inulin.

Inosite, C₆H₁₂O₆, 2H₂O, is not a carbohydrate, but, according to Hammersten, is an aromatic compound. Besides occurring in unripe fruits, it is found in green asparagus and beans.

Pectose is a substance the exact nature of which has not been fully determined, though it is thought to be a carbohydrate. It gives to unripe fruits and vegetables their peculiar hardness, and furnishes the basis for their gelatinous constituents. When the vegetable or fruit ripens, the insoluble pectose is then transformed by the action of acids and possibly of ferments into pectin, a vegetable jelly, which gives to fruit juice the property of gelatinizing when boiled.

Inulin, $(C_6H_{10}O_5)_n$, is a starch-like substance, occurring in the roots of chicory and dandelion, and in the tubers of the artichoke. It is a white, starch-like powder, slightly soluble in cold, and readily soluble in hot water, and converted into levulose by boiling with water, or by the action of acids.

METHODS OF PROXIMATE ANALYSIS.

Preparation of the Sample.—Cereals and dry leguminous foods are prepared for analysis by grinding in a coffee- or spice-mill to such a degree of fineness that the powder will pass through a sieve with 60 meshes to the inch. Green vegetables, beets, green peas, etc., are best reduced to suitable form for analysis by running through a domestic grinding-machine of the kind ordinarily employed in the kitchen for grinding and shredding meats and vegetables, being by this means reduced to a pulp of uniform consistency.

The following methods are based for the most part on those of the Association of Official Agricultural Chemists, employed for the analysis of foodstuffs with modifications.*

^{*} U. S. Dept. of Agric., Div. of Chem., Bul. 46 (rev.), and Bul. 107 (rev.).

Moisture.—Two grams of the substance are dried at 100° C. for five hours in a current of dry hydrogen, in a suitable drying oven. Results sufficiently accurate for most purposes may be secured by weighing the substance into a weighing-bottle or dish, and drying without hydrogen in an ordinary water oven.

Ash.—Two grams of the substance are burned in a platinum dish to whiteness at the lowest possible red heat.* If a white ash cannot be obtained in this manner, exhaust the charred mass with water, collect the insoluble residue on a filter, burn, add this ash to the residue from the evaporation of the aqueous extract, and heat the whole to a low redness till the ash is white or nearly so.

Ether Extract (Fat, etc.).—The residue from the determination of moisture is extracted for sixteen hours with anhydrous, alcohol-free ether in a continuous extractor. The extract is dried to constant weight, or the ether extract may be determined indirectly from the difference in weight of the dried substance before and after extraction, weighing it for convenience in the extraction tube.

Protein.—The total nitrogen is determined according to the Gunning or Kjeldahl method in the absence of nitrates, using I gram of the finely divided substance. The protein is calculated by multiplying the total nitrogen by the appropriate factor, which varies with the different cereals as follows: wheat, 5.70; rye, 5.62; oats, 6.31; corn, 6.39; and barley, 5.82. Ordinarily the conventional factor 6.25 is employed.

Crude Fiber (Cellulose, Lignin, etc.).†—The residue, after extraction for the determination of the ether extract, is transferred to a 500-cc. flask, with a mark showing 200 cc., and boiling 1.25% sulphuric acid is added to the mark. Heat at once to boiling, and boil gently for thirty minutes, shaking cautiously from time to time to prevent the material from crawling up on the sides of the flask. Filter through paper, and wash once with boiling water. Rinse the substance back into the same flask with 200 cc. of a boiling 1.25% solution of sodium hydroxide, free, or nearly so, from sodium carbonate, boil at once, and continue the boiling for thirty minutes in the same manner as directed above for the treatment with acid. Filter on a tared filter-paper, and wash with boiling water till the washings are neutral. Dry at 110° and weigh, after which incinerate completely. The loss of weight is crude fiber. A blank experiment should

^{*} Observe the precautions, indicated on page 134 about igniting cereals in platinum. † Modified from U. S. Dept. of Agric., Div. of Chem., Bul. 46.

be made on a second piece of filter-paper to show the loss occasioned by treatment with alkali, and the necessary correction should be made.

The filter used for the first filtration may be linen, one of the forms of glass wool or asbestos filters, or any other form that secures clear and reasonably rapid filtration. A gooch was originally prescribed for the final filtration, but with many substances is apt to clog. The solutions of sulphuric acid and sodium hydroxide are to be made up of the specified strength determined accurately by titration, and not merely from specific gravity.

Nitrogen-free Extract (Starch, Sugar, Gums, etc.).—Subtract the sum of the moisture, ash, ether extract, protein, and crude fiber, from 100.

Determination of Moisture in Grain.—Brown and Duvel Method.*

—This method was devised for purposes of inspection with the view

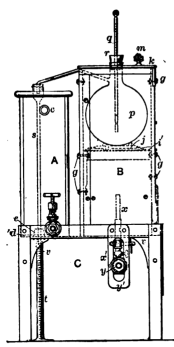


Fig. 61.—Brown and Duvel Apparatus for Determination of Water in Grain. End View.

of guarding against an excessive amount of moisture in corn, which causes deterioration through the growth of bacteria and moulds. The determinations are made on the whole grain in the apparatus shown in Fig. 61. This consists of a condenser-tank (A) and an evaporatingchamber (B) with a cover (k), the whole supported on a stand (C). Each of the flasks (b) rests on a flanged pipe-stem triangle, which in turn rests on a wire gauze. The apparatus is arranged for conducting six distillations at the same time.

Introduce into the distillation-flask (p) 100 cc. of a good grade hydrocarbon oil, and 100 grams of the grain (weighed on a torsion balance accurate to 0.03 gram), and close the neck of the flask with a rubber stopper carrying a thermometer (q), the bulb of which extends well into the mixture of oil and corn. Connect the side tube of the flask by means

of another cork with the condenser-tube (s), and heat with the Bunsen

burner until the thermometer registers 190° C., which requires from ten to fifteen minutes according to the amount of moisture present and the size of the flame. Turn off the flame, and allow to stand eight to ten minutes, or until the moisture ceases to drop from the condenser-tube into the graduate (t). The number of cc. in the graduate represents the percentage of moisture in the grain.

The results agree closely with those by drying to constant weight in a water oven at 100°.

The hydrocarbon oil should have a flash-point, in an open cup, of from 200° to 205° C. It is sold under the name of engine oil.

CARBOHYDRATES OF CEREALS AND VEGETABLES.

Classification.—As a rule the same carbohydrates are found in all cereals, being present, however, in varying proportions. By far the greater part of the carbohydrate content of cereals is starch, the other carbohydrates being comparatively small in amount, so that in rough work it is sometimes customary, though incorrect, to assume the entire amount of so-called "nitrogen-free extract" or carbohydrates (as determined by difference) to be starch.

The carbohydrates occurring in cereals may be classified as follows:

Principal carbohydrates of cereals:	Insoluble	
of cereals:	Soluble	Sucrose Dextrose Dextrin Raffinose (traces)

STARCH $(C_5H_{10}O_5)_n$.—Pure starch is a glistening, white, granular powder having a peculiar feeling when rubbed between the thumb and finger. It is a very hygroscopic, commercial starch containing about 18% of moisture. Starch is very widely distributed in the vegetable kingdom, occurring in almost every plant at some stage in its growth.

Starch is insoluble in cold water, alcohol, and ether; it is soluble in hot water, though not without change. By boiling with dilute acids, starch is first converted by hydrolysis into a mixture of dextrin and maltose, and finally by prolonged boiling into dextrose. Malt extract also hydrolizes starch in solution.

Detection.—Starch is best detected, when present to any appreciable extent in any mixture, by boiling a portion of the sample in water, cooling, and applying a solution of iodine. A characteristic blue color is produced if starch is present. Very small amounts of starch are best iden-

tified in powdered mixtures by applying a drop of a solution of iodine to the dry powder on a microscope slide, or, better, to the powder previously rubbed out with water on a slide under a cover-glass; the starch granules, if present, will be colored intensely blue by the iodine, and are at once rendered apparent when viewed through the microscope.

Though the cereal and vegetable starches, whatever their origin, are identical chemically, the various starch granules have certain characteristics, when viewed under the microscope, that render their identification easy in most cases. A knowledge of the microscopical appearance of the common vegetable starches is of the utmost importance to the public analyst, who frequently finds them as adulterants of various foods, such as coffee, cocoa, spices, etc. For microscopical examination, powdered samples should be ground fine enough to pass through a 60 or 80 mesh sieve.

Classification.—The microscopical appearance of the starch granules of various grains and vegetables differ in form, size, and often in their manner of grouping. Thus, at the outset, the common starches may be divided as to the microscopical form of their granules into three classes, viz., circular, irregularly oval, and polygonal. To the first class, in which the starch granule has in general the circular disk form, belong rye, wheat, and barley. Representing the second or irregularly elliptical class are the pea, bean, potato, and arrowroot. In the third, or polygonal class, should be included corn, oats, buckwheat, and rice. In thus characterizing the distinguishing forms as circular, oval, and polygonal, it should be borne in mind that while the tendency of the most typical starch granules in each class, when viewed in normal position, is toward the circular, the oval, or the polygonal as the case may be, it is not by any means true that all or even most of the granules in any one instance perfectly conform to one of these shapes throughout.' Thus, circular wheat granules, when viewed edgewise, will appear elliptical, and are often distorted in shape, especially when roasted; and polygonal buckwheat granules may in many instances have such obtuse angles as to appear circular. It is the general trend of all the starches toward one or another of these shapes that suggests the classification.

The identification of the various starches morphologically is indeed the most natural and ready method. Not only the character of the starch, but also its approximate amount, when present in mixtures, can in many instances be ascertained by a careful examination with the microscope. The analyst should be provided with samples of starches of known purity conveniently at hand, and in all doubtful cases these should be referred to for comparison.

Wheat Starch (Fig. 152, Pl. VIII).—This starch is frequently present in adulterated pepper, mustard, ginger, cocoa, coffee, and other foods. Its granules occur for the most part in two sizes, of which the larger are circular disks, varying from 0.021 mm. to 0.041 mm., or rarely 0.050 mm., in diameter, while the smaller are rounded or polygonal, averaging about 0.005 mm. in diameter. The smaller granules are grouped irregularly in and around the larger, there being six to ten of the former to one of the latter. The larger granules are, however, the most distinctly characteristic, and are usually readily recognized in a mixture, not only by their shape, but by reason of the concentric rings with which they are provided, and which are generally but not always apparent.

Barley Starch (Fig. 124, Pl. I).—This much resembles wheat, in that it has two sizes of granules, but both sizes are respectively smaller than those of wheat, though present in about the same proportion. The larger circular disk-like granules vary from 0.013 mm. to 0.035 mm. in diameter, while the smaller average 0.003 mm. The concentric rings are less apparent in the barley than in the wheat.

Rye Starch (Fig. 148, Pl. VII) has also two sizes of granules, but the larger vary from 0.025 mm. to over 0.05 mm. in diameter, and are considerably larger than the corresponding wheat granules. The smaller granules average about 0.004 mm. in diameter. As in the case of wheat and barley, the larger granules are circular disks, while the smaller are rounded or polygonal. The concentric rings are usually indistinct in the large granules, and many of these show cross-shaped rifts in the center.

Corn Starch (Fig. 133, Pl. IV).—This starch is a common adulterant of spices, cocoa, and other foods. It is placed in a series of four cereal starches whose granules are polygonal, and all of which show more or less tendency to arrange themselves in close contact side by side in masses suggestive of a tessellated or mosaic floor. Arranged in order of the size of their grains, these starches are: Corn, buckwheat, oats, and rice. Corn starch granules tend toward the hexagonal in shape, varying from 0.007 mm. to 0.035 mm. in diameter, and having very marked rifted hila. They are most readily recognized in any mixture, and from their size are readily distinguishable from the other polygonal starches, which never reach 0.017 mm. in diameter.

Buckwheat Starch (Fig. 128, Pl. II, and Fig. 129, Pl. III).—This is a very common adulterant of many spices, especially pepper, which, as

shown in Fig. 256, Pl. XXXIV, it much resembles in the manner in which its masses of granules group themselves, conforming to the shape of the cells. The individual granules are commonly 0.006 mm. to 0.012 mm. in diameter. Curious rod-shaped aggregates of two to four individuals are of frequent occurrence.

Oat Starch (Fig. 139, Pl. V).—The granules of this starch vary from 0.002 mm. to 0.012 mm. in diameter, and are polygonal, or less often rounded or spindle-shaped in form. They have no rings or hila, and arrange themselves in rounded aggregates of from two to many granules that at first sight might be mistaken for large grains; careful examination, however, shows the dividing lines.

Rice Starch (Fig. 143, Pl. VI).—The granules of rice starch resemble closely those of oats both in form and size, but spindle-shaped forms are not present. As in the case of oats, the granules are often united to form rounded aggregates.

Starches of the Pea and Bean.—The starches of these legumes much resemble each other, and are with difficultly distinguished one from the other (see Fig. 164, Pl. XI, and Fig. 154, Pl. IX). The granules are more nearly oval than most other starches, and have both concentric rings and elongated hila. The granules of the pea show a less distinct hilum than those of the bean, and some of them are irregularly swollen. Both peas and beans roasted are commonly used as adulterants of coffee.

Arrowroot.—There are many varieties of arrowroot, including Jamaica, Bermuda, East Indian, Australian, and others, all having certain variations in form and size, but resembling each other in a general way. Fig. 167, Pl. XII, shows the Bermuda arrowroot, the granules of which are somewhat egg-shaped, being usually smaller at one end than the other, and having rifted hila near the small end.

Potato Starch (Fig. 165, Pl. XII).—This starch has large, irregularly oval granules, with very apparent hila situated eccentrically near one end, and with rings around the hilum. The granules are about 0.07 mm. in large diameter. Fig. 134, Pl. IV, and Fig. 166, Pl. XII, show corn and potato starch when viewed with polarized light with crossed Nicol prisms, the specimens being mounted in Canada balsam.

Tapioca Starch.—The granules of this starch, as shown in Fig. 168, Pl. XII, are more uniform in size throughout than those already described, averaging about 0.018 mm. in diameter, and being quite smoothly circular, without concentric rings, but having a distinctly dotted hilum in

the center. Many of the grains are cup-shaped, as if a segment of the circle had been removed.

Sago Starch (Fig. 172, Pl. XIII).—The granules of sago starch vary much in size, and might be called irregularly ellipsoidal in shape, being provided with numerous protuberances. Some of them have indistinct concentric rings, and in some, but not all, a hilum is apparent, usually near one end of the granule.

Microscopical Appearances of Starches with Polarized Light.—With polarized light starch granules show dark crosses, the point of intersection being at the hilum (Fig. 166, Pl. XII). These crosses vary in distinctness with the variety. Certain of the starches show a play of colors with polarized light and a selenite plate, especially those whose granules have some sort of hilum. This is particularly striking in such starches as corn, tapioca, potato, and arrowroot. Blyth has made the phenomenon a means of classification of the starches, but the writer considers their appearance with ordinary light sufficient for identification. Canada balsam is the best mountant for examination in polarized light.

Estimation of Starch.—Direct Acid Conversion.—By this method the hemicellulose, if present, or such of the carbohydrates as are capable of being converted to sugar, are reckoned in with the starch. Where little or none of the insoluble carbohydrates other than starch are present, as for instance in the case of commercial starches, this method is sufficiently accurate.

Exhaust 3 grams of the finely divided substance on a fine but rapidly acting filter with ether by washing with 5 successive portions of 10 cc. each, and wash the residue first with 150 cc. of 10% alcohol and then with a little strong alcohol. Transfer by washing to a flask with 200 cc. of water and 20 cc. of hydrochloric acid (specific gravity 1.125), connect with a reflux condenser, and heat the flask in boiling water for 2½ hours. Cool, and carefully neutralize with sodium hydroxide, clarifying if necessary with alumina cream. Mix well, make up the volume to 500 cc., filter, and determine the dextrose in an aliquot part of the filtrate by any of the methods for dextrose. Convert dextrose to starch by the factor 0.9.

Diastase Method.—By this method the hemicellulose is not converted, only the starch being acted upon. Hence for exact work in the presence of other insoluble carbohydrates this method is to be recommended. Under the action of diastase, starch is first converted into

maltose and dextrin, and finally into dextrose, in somewhat the following manner:

$$\begin{split} & \text{I2C}_6 \text{H}_{10} \text{O}_5 + 4 \text{H}_2 \text{O} = 4 \text{C}_{12} \text{H}_{22} \text{O}_{11} + 2 \text{C}_{12} \text{H}_{20} \text{O}_{10} \\ & \text{Starch} \end{split}$$

$$& \text{C}_{12} \text{H}_{22} \text{O}_{11} + \text{H}_2 \text{O} = 2 \text{C}_6 \text{H}_{12} \text{O}_6 \\ & \text{Maltose} \end{split}$$

$$& \text{C}_{12} \text{H}_{20} \text{O}_{10} + 2 \text{H}_2 \text{O} = 2 \text{C}_6 \text{H}_{12} \text{O}_6 \\ & \text{Maltose} \end{split}$$

Exhaust 3 grams of the finely divided substance with ether and alcohol as in the acid conversion method, wash the residue into a beaker with 100 cc. of water and boil directly for 30 minutes, stirring constantly and restoring the water lost on evaporation. Cool to 55° C., add 20 cc. of malt extract (prepared as below), and maintain at this temperature for one hour with occasional stirring. Boil a second time for 15 minutes. cool again to 55° C., and digest once more with 10 cc. of additional malt The treatment with malt converts the starch into dextrin and maltose. Heat to boiling a third time, cool and make up to 250 cc. Filter, transfer 200 cc. of the filtrate to a 500 cc. flask, and add 20 cc. of hydrochloric acid (specific gravity 1.125), connect with a reflux condenser and heat in a boiling-water bath for two and one-half hours. by which process the dextrin and maltose are converted into dextrose. Cool, neutralize carefully with sodium hydroxide (avoiding an excess), clarify if necessary with 10 to 20 cc. of alumina cream (p. 587), and make up to 500 cc. Mix well, pour through a dry filter, and determine the dextrose in an aliquot part of the filtrate, using the factor o.g for converting dextrose to starch. Correct for the copper reducing power of the malt extract, as below.

Preparation of Malt Extract.—Dry malted barley can be readily obtained from any brewery. Treat 15 to 20 grams of freshly pulverized malt for several hours with 100 cc. of water, shaking occasionally. Filter the solution, and add two or three drops of chloroform to prevent the growth of fungi. Determine the amount of dextrose in a given quantity of the malt extract, after boiling with acid, etc., as in the starch determination, and make the proper correction.

Use of "Animal Diastase."—Pancreatin and similar powdered preparations, such as "vera diastase" and "panase," obtained from the pancreas of cattle or hogs, are convenient for use as starch-converting reagents instead of malt extract, and are preferable to the latter in that, as a rule, they possess no copper-reducing ingredient and hence need no correction.

If of the strength of U. S. P. pancreatin, which should convert at least twenty-five times its weight of starch, use instead of the malt extract the same amount, viz., 20 cc., of a 0.5% aqueous solution of the powdered substance in starch determinations as above described.

Cellulose forms the framework, or skeleton, of all vegetable organisms, being, next to water, the most abundant substance in the vegetable kingdom.

Pure cellulose is white, translucent, and of fibrous or silky texture. It is insoluble in water, alcohol, and ether, but dissolves readily in an ammoniacal solution of cupric hydroxide known as Schweitzer's Reagent * or "cuprammonia."

Cellulose turns violet when treated with chloriodide of zinc, and blue when treated with sulphuric acid and iodine in potassium iodide (p. 91).

The "crude fiber" as determined in foods, being the portion that resists the action of hot dilute acid and alkali, is composed largely of cellulose.

The Pentosans are of comparatively small importance, and have been little studied. They are amorphous in character, insoluble in water, but soluble in dilute alkali, and are capable of conversion by boiling with dilute acids into so-called pentose sugars, the best known of which are xylose and arabinose, corresponding to the pentosans xylan and araban respectively. Strictly speaking the term "hemicellulose" is the more appropriate generic term to apply to the insoluble carbohydrate bodies which are capable of hydrolysis by acids to sugars, inasmuch as there are insoluble bodies besides the pentosans that may thus be converted into sugar, such as the hexosans, hydrolyzed by acid to hexose sugars, mannose, galactose, etc. The term "wood gum" is also used synonymously with hemicellulose. Since the greater portion of these insoluble hydrolizable carbohydrates are pentosans, it is simpler to calculate them all as such.

Determination of Pentosans.—Pentosans are determined either by hydrolyzing to reducing sugar, and estimating the latter as described on page 296 (Stone's method) or by calculation from the furfural† yielded

^{*} Prepared as directed on page 93.

[†] Furfural or furfuraldehyde (C₄H₄O₂) is the aldehyde of pyromucic acid. It is a color-less liquid, having an odor suggestive of cassia. Its boiling-point is 162° and its specific gravity 1.164. It is sparingly soluble in water and readily soluble in alcohol. Nearly half the tissue of ordinary bran, exclusive of proteins and starch, yields furfural on distillation with acid.

by them on distillation of the sample with hydrochloric acid, as carried out in the provisional method of the A. O. A. C.* as follows:

Three grams of the material are placed in a flask, together with 100 cc. of 12% hydrochloric acid (specific gravity 1.06) and several pieces of recently heated pumice stone. The flask, placed upon wire gauze, is connected with a condenser, and heat applied, rather gently at first, using a gauze top to distribute the flame, and so regulated as to distill over 30 cc. in about ten minutes, passing the distillate through a small filter. The 30 cc. driven over are replaced by a like quantity of the dilute acid, and the process continued until the distillate amounts to 360 cc. To the completed distillate is gradually added a quantity of phloroglucinol (free from diresorcin) dissolved in 12% of hydrochloric acid, and the resulting mixture is thoroughly stirred. The amount of phloroglucinol used should be about double that of the furfural expected. The solution first turns yellow, then green, and very soon an amorphous greenish precipitate appears, which grows rapidly darker, till it finally becomes almost black. The solution is made up to 400 cc. with 12% hydrochloric acid and allowed to stand over night.

The amorphous, black precipitate, a condensation product the exact composition of which is unknown, is filtered into a tared gooch through an asbestos felt, washed with 150 cc. of water in such a way that the precipitate is kept covered with liquid until the last portion has passed through the filter, dried to constant weight by heating from three to four hours at 100°, cooled in a weighing-bottle, and weighed, the increase in weight being reckoned as phloroglucide. To calculate the furfural, pentoses, and pentosans from the phloroglucide, use Kröber's formulæ as follows:

(a) For weight of phloroglucide "a" under 0.03 gram.

Furfural = $(a+0.0052) \times 0.5170$. Pentoses = $(a+0.0052) \times 1.0170$. Pentosans = $(a+0.0052) \times 0.8949$.

^{*} U. S. Dept. of Agric., Bur. of Chem., Bul. 107 (rev.), p. 54.

`:.

(b) For weight of phloroglucide "a" over 0.300 gram.

Furfural =
$$(a+0.0052) \times 0.5180$$
.
Pentoses = $(a+0.0052) \times 1.0026$.
Pentosans = $(a+0.0052) \times 0.8824$.

For weight of phloroglucide "a" from 0.03 to 0.300 gram use Kröber's table (pp. 288-294) to calculate the weight of pentoses (arabinose, xylose), and pentosans (araban, xylan).

The reactions that take place are thought to be somewhat as follows:

$$C_5H_8O_4+H_2O=C_5H_{10}O_5$$
.
Pentosan

$$C_5H_{10}O_5 = C_5H_4O_2 + 3H_2O_5$$

Pentose Furfural

$$2C_5H_4O_2+C_6H_6O_3=C_{16}H_{12}O_6+H_2O_6$$

Furfural Phloroglucinol Phloroglucide

The theoretical yield of phloroglucide should be 2.22 parts to one of furfural, but in practice this is never obtained. The varying factors for calculation as above given are based on experiment.

The phloroglucinol used should be free from diresorcin. To test for the latter, dissolve the reagent in acetic anhydride, heat nearly to boiling, and add a few drops of concentrated sulphuric acid. If more than a faint violet color is produced, the phloroglucinol should be purified as follows:

Heat in a beaker about 300 cc. of hydrochloric acid (sp. gr. 1.06) and 11 grams of commercial phloroglucinol, added in small quantities at a time, stirring constantly until it has almost dissolved. Some impurities may resist solution, but it is unnecessary to dissolve them. Pour the hot solution into a sufficient quantity of the same hydrochloric acid (cold) to make the volume 1500 cc. Allow it to stand at least overnight—better several days—to allow the diresorcin to crystallize out, and filter immediately before using. The solution may turn yellow, but this does not interfere with its usefulness. In using it, add the volume containing the required amount to the distillate.

KRÖBER'S TABLE FOR DETERMINATION OF PENTOSES AND PENTOSANS FROM PHLOROGLUCID.

Phloroglucid	Furfural.	Arabinose.	Araban.	Xylose.	Xylan.	Pentose.	Pentosan
0.030	0.0182	0.0391	0.0344	0.0324	0.0285	0.0358	0.0315
.031	.0188	.0402	-0354	.0333	.0293	.0368	.0324
.032	.0193	.0413	.0363	.0342	.0301	.0378	.0333
.033	.0198	.0424	-0373	.0352	.0309	.0388	.0341
.034	.0203	.0435	.0383	.0361	.0317	.0398	.0350
-035	.0209	.0446	.0393	.0370	.0326	.0408	.0359
.036	.0214	.0457	.0402	-0379	-0334	.0418	.0368
.037	.0219	.0468	.0412	.0388	.0342	.0428	-0377
.038	.0224	-0479	.0422	.0398	.0350	.0439	.0386
.039	.0229	.0490	.0431	.0407	.0358	-0449	-0395
.040	.0235	.0501	.0441	.0416	.0366	.0459	.0404
.041	.0240	.0512	.0451	.0425	.0374	.0469	.0413
-042	.0245	.0523	.0460	.0434	.0382	.0479	.0422
-043	.0250	-0534	.0470	-0443	.0390	.0489	.0431
.044	-0255	-0545	.0480	.0452	.0398	-0499	.0440
.045	.0260	.0556	.0490	.0462	.0406	.0509	.0448
.046	.0266	.0567	.0499	.0471	.0414	.0519	-0457
-047	.0271	.0578	.0509	-0480	.0422	.0529	.0466
.048	.0276	.0589	.0519	.0489	.0430	.0539	.0475
.049	.0281	.0600	.0528	-0498	.0438	-0549	.0484
, • 050	.0286	.0611	.0538	-0507	.0446	.0559	.0492
.051	.0292	.0622	.0548	-0516	-0454	.0569	.0501
.052	.0297	-0633	-0557	.0525	.0462	.0579	.0510
•053	.0302	.0644	.0567	-0534	.0470	.0589	.0519
.054	.0307	.0655	.0576	.0543	.0478	.0599	.0528
-055	.0312	.0666	.0586	-0553	.0486	.0610	.0537
.056	.0318	.0677	.0596	.0562	.0494	.0620	.0546
.057	.0323	.0688	.0605	.0571	.0502	.0630	.0555
.058	.0328	.0699	.0615	.0580	.0510	.0640	.0564
.059	.0333	.0710	.0624	.0589	.0518	.0650	-0573
.060	.0338	.0721	.0634	.0598	.0526	.0660	.0581
.061	-0344	.0732	.0644	.0607	-0534	.0670	.0590
.062	-0349	-0743	.0653	.0616	.0542	.0680	.0599
.063	.0354	-0754	.0663	.0626	.0550	.0690	.0608
.064	.0359	-0765	.0673	.0635	.0558	-0700	.0617
.065	.0364	.0776	.0683	.0644	.0567	.0710	.0625
.066	.0370	-0787	.0692	.0653	-0575	.0720	.0634
.067	-0375	.0798	.0702	.0662	.0583	.0730	.0643
.068	.0380	.0809	.0712	.0672	.0591	.0741	.0652
.069	.0385	.0820	.0721	.0681	.0599	.0751	.0661

KRÖBER'S TABLE FOR DETERMINATION OF PENTOSES AND PENTOSANS FROM PHLOROGLUCID—Continued.

Phloroglucid	Purfural.	Arabinose.	Araban.	Xylose.	Xylan.	Pentose.	Pentosan.
0.070	0.0390	0.0831	0.0731	0.0690	0.0607	0.0761	0.0670
.071	.0396	.0842	.0741	.0699	.0615	.0771	.0679
.072	.0401	.0853	.0750	.0708	.0623	.0781	.0688
.073	.0406	.0864	.0760	.0717	.0631	.0791	.0697
.074	.0411	.0875	.0770	.0726	.0639	10801	.0706
•075	.0416	.0886	.0780	.0736	.0647	.0811	.0714
.076	.0422	.0897	.0789	.0745	.0655	.0821	.0722
-077	.0427	.0908	.0799	-0754	.0663	.0831	.0731
.078	.0432	.0919	.0809	.0763	.0671	.0841	.0740
.079	-0437	.0930	.0818	.0772	.0679	.0851	-0749
.080	.0442	.0941	.0828	.0781	.0687	.0861	.0758
.081	.0448	.0952	.0838	.0790	.0695	.0871	.0767
.082	.0453	.0963	.0847	-0799	.0703	.0881	.0776
.083	.0458	.0974	.0857	.0808	.0711	.0891	.0785
.084	.0463	.0985	.0867	.0817	.0719	10901	-0794
.085	.0468	.0996	.0877	.0827	.0727	.0912	. 0803
.086	.0474	.1007	.0886	.0836	-0735	.0922	.0812
.087	.0479	.1018	.0896	.0845	.0743	.0932	.0821
.088	.0484	.1029	.0906	.0854	.0751	.0942	.0830
.089	.0489	.1040	.0915	.0863	-0759	.0952	.0838
.090	.0494	.1051	.0925	.0872	.0767	.0962	.0847
.091	.0499	.1062	-0935	.0881	-0775	.0972	.0856
.092	.0505	.1073	.0944	.0890	-0783	.0982	.0865
•093	.0510	.1084	.0954	.0900	.0791	.0992	.0874
.094	.0515	.1095	.0964	.0909	.0800	.1002	.0883
.095	.0520	.1106	-0974	.0918	.0808	.1012	.0891
.096	.0525	.1117	.0983	.0927	.0816	.1022	.0899
-097	.0531	.1128	.0993	.0936	.0824	.1032	.0908
.098	.0536	.1139	.1003	.0946	.0832	.1043	.0917
.099	.0541	.1150	.1012	-0955	.0840	.1053	.0926
.100	.0546	.1161	.1022	.0964	.0848	.1063	.0935
.101	.0551	.1171	.1032	.0973	.0856	.1073	•0944
.102	.0557	.1182	.1041	.0982	.0864	.1083	.0953
.103	.0562	.1193	.1051	.0991	.0872	.1093	.0962
.104	.0567	.1204	.1060	.1000	.0880	.1103	.0971
.105	.0572	.1215	.1070	.1010	.0888	.1113	.0976
.106	.0577	.1226	.1080	.1019	.0896	.1123	.0988
.107	.0582	.1237	.1089	.1028	.0904	.1133	-0997
.108	.0588	.1248	.1099	.1037	.0912	.1143	.1006
.109	.0593	.1259	.1108	.1046	.0920	.1153	.1015

KRÖBER'S TABLE FOR DETERMINATION OF PENTOSES AND PENTOSANS FROM PHLOROGLUCID—(Continued).

Phloroglucid	Furfural.	Arabinose.	Araban.	Xylose.	Xylan.	Pentose.	Pentosan.
0.110	0.0598	0.1270	0.1118	0.1055	0.0928	0.1163	0.1023
.111	.0603	.1281	.1128	.1064	.0936	-1173	.1032
.112	.0608	.1292	.1137	.1073	.0944	.1183	.1041
.113	.0614	.1303	.1147	.1082	.0952	.1193	1050
.114	.0619	.1314	.1156	.1091	.0960	.1203	. 1059
.115	.0624	.1325	.1166	.1101	.0968	.1213	.1067
.116	.0629	.1336	.1176	.1110	.0976	.1223	.1076
.117	.0634	-1347	.1185	.1119	.0984	.1233	.1085
.118	.0640	.1358	.1195	.1128	.0992	.1243	.1094
.119	.0645	.1369	. 1 204	.1137	.1000	.1253	.1103
. 1 20	.0650	.1380	.1214	.1146	.1008	.1263	.1111
.121	.0655	.1391	. 1 2 2 4	.1155	.1016	.1273	.1120
.122	.0660	.1402	.1233	.1164	.1024	.1283	.1129
.123	.0665	.1413	-1243	.1173	.1032	.1293	.1138
.124	-0671	.1424	.1253	.1182	.1040	.1303	-1147
.125	.0676	.1435	. 1 263	.1192	. 1049	.1314	.1156
.126	.0681	.1446	.1272	.1201	.1057	.1324	.1165
.127	.0686	-1457	. 1282	.1210	.1065	-1334	-1174
. 1 28	.0691	.1468	. 1 292	.1219	.1073	.1044	.1183
.129	.0697	-1479	.1301	.1228	.1081	-1354	.1192
.130	.0702	.1490	.1311	.1237	. 1089	.1364	.1201
.131	-0707	.1501	.1321	.1246	.1097	.1374	.1210
.132	.0712	.1512	.1330	-1255	.1105	.1384	.1219
-133	.0717	-1523	.1340	.1264	.1113	.1394	.1227
-134	.0723	-1534	.1350	-1273	.1121	.1404	.1236
.135	.0728	.1545	.1360	. 1 283	.1129	.1414	. 1 244
.136	.0733	.1556	.1369	.1292	-1137	.1424	.1253
-137	.0738	.1567	.1379	.1301	.1145	-1434	.1262
.138	.0743	.1578	.1389	.1310	.1153	-1444	.1271
-139	.0748	.1589	.1398	.1319	.1161	-1454	. 1 280
.140	.0754	.1600	. 1408	.1328	.1169	.1464	. 1 288
.141	.0759	.1611	.1418	-1337	.1177	-1474	.1297
.142	.0764	.1622	.1427	.1346	.1185	.1484	.1306
-143	.0769	.1633	-1437	1355	.1193	.1494	.1315
-144	-0774	.1644	-1447	.1364	.1201	.1504	.1324
.145	.0780	.1655	.1457	.1374	.1209	.1515	-1333
.146	.0785	.1666	_1466	.1383	.1217	.1525	.1342
-147	.0790	.1677	.1476	.1392	.1225	.1535	.1351
.148	•0795	.1688	. 1486	.1401	.1233	.1545	.1360
.149	.0800	.1699	. 1495	.1410	.1241	-1555	.1369

KRÖBER'S TABLE FOR DETERMINATION OF PENTOSES AND PENTOSANS FROM PHLOROGLUCID-(Continued).

Phloroglucid	Furfural.	Arabinose,	Araban.	Xylose.	Kylan.	Pentose.	8 Pentosan.
0.150	0.0805	0.1710	0.1505	0.1419	0.1249	0.1565	0.1377
.151	.0811	.1721	.1515	.1428	.1257	-1575	.1386
.152	.0816	.1732	. 1524	-1437	.1265	. 1585	.1395
.153	.0821	-1743	.1534	.1446	.1273	-1595	.1404
-154	.0826	-1754	-1544	-1455	.1281	.1605	.1413
.155	-0831	.1765	-1554	.1465	.1289	.1615	.1421
.156	-0837	-1776	.1563	-1474	.1297	. 1625	.1430
-I57	-0842	.1787	-1573	.1483	.1305	. 1635	.1439
.158	-0847	.1798	. 1583	.1492	.1313	-1645	.1448
-159	.0852	.1809	.1592	.1501	.1321	.1655	-1457
.160	.0857	.1820	. 1602	.1510	.1329	.1665	.1465
.161	.0863	.1831	. 1612	.1519	-1337	.1675	.1474
.162	.0868	.1842	.1621	.1528	.1345	.1685	.1483
.163	.0873	.1853	.1631	-1537	1353	.1695	.1492
.164	-0878	.1864	.1640	.1546	.1361	.1705	.1501
.165	.0883	-1875	. 1650	.1556	.1369	.1716	.1510
.166	.0888	.1886	.1660	.1565	-1377	.1726	.1519
.167	.0894	. 1897	. 1669	-1574	.1385	.1736	.1528
.168	.0899	.1908	.1679	.1583	-1393	.1746	.1537
.169	.0904	.1919	. 1688	.1592	.1401	.1756	.1546
.170	.0909	.1930	1698	.1601	.1409	1766	1554
-171	.0914	.1941	. 1708	.1610	-1417	.1776	.1563
.172	.0920	-1952	.1717	.1619	.1425	.1786	.1572
-173	.0925	-1963	.1727	.1628	-1433	.1796	.1581
-174	.0930	-1974	.1736	.1637	-1441	.1806	.1590
-175	-0935	.1985	.1746	.1647	-1449	.1816	.1598
.176	-0940	.1996	.1756	.1656	-1457	.1826	.1607
-177	.0946	.2007	. 1765	.1665	.1465	.1836	.1616
.178	.0951	.2018	-1775	.1674	-1473	.1846	.1625
-179	.0956	.2029	.1784	.1683	.1481	.1856	.1634
.180	.0961	.2039	1794	.1692	.1489	.1866	.1642
.181	.0966	-2050	. 1804	.1701	-1497	.1876	.1651
.182	.0971	. 2061	.1813	.1710	.1505	.1886	.1660
.183	.0977	.2072	. 1823	.1719	.1513	.1896	.1669
.184	.0982	. 2082	. 1832	.1728	.1521	.1906	.1678
.185	.0987	. 2093	.1842	.1738	.1529	.1916	.1686
. 186	.0992	.2104	.1851	-1747	.1537	.1926	.1695
.187	-0997	.2115	. 1861	.1756	-1545	.1936	.1704
.188	.1003	.2126	.1870	.1765	-1553	.1946	.1712
. 189	.1008	.2136	.1880	-1774	.1561	.1955	.1721

KRÖBER'S TABLE FOR DETERMINATION OF PENTOSES AND PENTOSANS FROM PHLOROGLUCID—(Continued).

1 Phloroglucid	Purfural.	Arabinose.	Araban.	Xylose.	Kylan.	7 Pentose.	8 Pentosan.
0.190	0.1013	0.2147	0.1889	0.1783	0.1569	0.1965	0.1729
.191	. 1018	.2158	.1899	.1792	-1577	-1975	.1738
.192	.1023	.2168	.1908	.1801	.1585	.1985	-1747
.193	.1028	.2179	.1918	.1810	. 1593	.1995	.1756
-194	.1034	.2190	-1927	.1819	.1601	.2005	.1764
-195	.1039	.2201	-1937	.1829	.1609	.2015	-1773
.196	.1044	.2212	.1946	.1838	.1617	.2025	.1782
-197	.1049	. 2222	.1956	.1847	.1625	. 2035	.1791
-198	.1054	-2233	.1965	.1856	.1633	. 2045	.1800
-199	.1059	.2244	-1975	.1865	.1641	. 2055	.1808
.200	.1065	.2255	.1984	.1874	.1649	. 2065	.1817
.201	.1070	.2266	.1994	.1883	.1657	. 2075	. 1826
.202	. 1075	.2276	.2003	.1892	.1665	.2085	.1835
. 203	.1080	.2287	.2013	.1901	.1673	. 2095	.1844
. 204	.1085	.2298	.2022	.1910	.1681	.2105	.1853
. 205	.1090	.2309	.2032	.1920	.1689	.2115	.1861
. 206	.1096	.2320	.2041	.1929	.1697	.2125	.1869
- 207	.1101	.2330	.2051	.1938	.1705	.2134	. 1878
_ 208	.1106	.2341	.2060	-1947	.1713	.2144	.1887
. 209	.1111	.2352	. 2069	.1956	.1721	.2154	.1896
.210	.1116	.2363	.2079	.1965	.1729	.2164	.1904
- 2I I	.1121	-2374	. 2089	-1975	-1737	-2174	.1913
.212	.1127	.2384	.2098	.1984	-1745	.2184	.1922
.213	.1132	-2395	.2108	.1993	-1753	.2194	.1931
-214	.1137	.2406	.2117	.2002	.1761	. 2204	.1940
.215	.1142	.2417	.2127	.2011	.1770	.2214	.1948
-216	.1147	.2428	.2136	.2020	.1778	. 2224	.1957
.217	.1152	.2438	.2146	.2029	.1786	.2234	.1966
.218	.1158	-2449	.2155	.2038	.1794	.2244	.1974
~219	.1163	.2460	.2165	.2047	.1802	. 2254	.1983
.220	.1168	.2471	.2174	.2057	.1810	. 2264	.1992
.221	.1173	.2482	.2184	. 2066	.1818	-2274	.2001
.222	.1178	- 2492	.2193	-2075	.1826	.2284	. 2010
.223	.1183	.2503	.2203	.2084	.1834	.2294	.2019
. 224	.1189	.2514	.2212	.2093	.1842	.2304	.2028
.225	.1194	.2525	.2222	.2102	.1850	.2314	2037
226	.1199	.2536	.2232	.2111	.1858	.2324	. 2046
-227	.1204	.2546	.2241	.2121	.1866	•2334	.2054
228	.1209	-2557	.2251	.2130	.1874	-2344	.2063
.229	.1214	.2568	.2260	.2139	.1882	-2354	.2072

KRÖBER'S TABLE FOR DETERMINATION OF PENTOSES AND PENTOSANS FROM PHLOROGLUCID—(Continued).

Phloroglucid	Purfural.	Arabinose.	Araban.	Xylose.	6 Xylan.	Pentose.	Pentosan
0.230	0.1220	0.2579	0.2270	0.2148	0.1890	0.2364	0.2081
.231	.1225	.2590	. 2280	.2157	.1898	-2374	. 2089
.232	.1230	. 2600	. 2289	.2166	.1906	.2383	. 2097
-233	.1235	.2611	. 2299	.2175	.1914	.2393	.2106
-234	.1240	. 2622	.2308	.2184	.1922	.2403	.2115
-235	.1245	. 2633	.2318	.2193	.1930	.2413	. 21 24
.236	.1251	.2644	. 2327	.2202	.1938	.2423	.2132
-237	.1256	.2654	-2337	.2211	.1946	-2433	.2141
.238	.1261	.2665	. 2346	.2220	1954	•2443	.2150
•239	. 1 266	. 2676	.2356	.2229	.1962	-2453	.2159
.240	.1271	. 2687	. 2365	.2239	.1970	.2463	.2168
.241	.1276	. 2698	-2375	.2248	.1978	•2473	.2176
. 242	.1281	.2708	. 2384	.2257	.1986	.2483	.2185
-243	. 1 287	.2719	- 2394	.2266	-1994	•2493	.2194
.244	.1292	.2730	.2403	.2275	.2002	.2503	.2203
.245	. 1 297	.2741	.2413	. 2284	. 2010	.2513	.2212
.246	.1302	.2752	.2422	.2293	.2018	-2523	. 2220
.247	. 1 307	.2762	.2432	.2302	.2026	·2533	. 2229
.248	.1312	-2773	. 2441	.2311	.2034	-2543	.2238
-249	1318	.2784	.2451	.2320	.2042	-2553	.2247
. 250	.1323	.2795	. 2460	.2330	. 2050	.2563	.2256
.251	.1328	.2806	. 2470	•2339	.2058	•2573	.2264
. 252	•1333	.2816	-2479	. 2348	. 2066	.2582	.2272
-253	.1338	. 2827	.2489	-2357	.2074	.2592	.2281
.254	.1343	. 2838	. 2498	.2366	.2082	. 2602	.2290
·255	.1349	. 2849	.2508	-2375	.2090	.2612	.2299
.256	-1354	. 2860	.2517	.2384	.2098	.2622	.2307
-257	.1359	. 2870	.2526	•2393	.2106	.2632	.2316
.258	.1364	. 2881	.2536	.2402	.2114	. 2642 . 2652	.2325
•259	.1369	.2092	-2545	.2411	.2122	.2052	-2334
.260	.1374	. 2903	.2555	.2420	.2130	.2662	.2343
.261	.1380	.2914	.2565	.2429	.2138	. 2672	. 2351
.262	.1385	. 2924	.2574	.2438	.2146	.2681	. 2359
.263	.1390	.2935	.2584	.2447	.2154	.2691	.2368
. 264	.1395	. 2946	.2593	. 2456	.2102	. 2701	•2377
.265	.1400	.2957	2603	. 2465	.2170	.2711	. 2385
. 266	.1405	. 2968	.2612	-2474	.2178	.2721	2394
. 267	.1411	.2978	. 2622	. 2483	.2186	.2731	. 2403
. 268	.1416	.2989	.2631	.2492	.2194	.2741	.2412
.269	.1421	.3000	.2641	.2502	.2202	.2751	.2421

KRÖBER'S TABLE FOR DETERMINATION OF PENTOSES AND PENTOSANS FROM PHLOROGLUCID—(Concluded).

Phloroglucid	Furfural.	Arabinose.	Araban.	Xy lose.	6 Xylan.	Pentose.	8 Pentosan.
0.270	0.1426	0.3011	0.2650	0.2511	0.2210	0.2761	0.2429
-271	-1431	.3022	.2660	.2520	.2218	.2771	. 2438
-272	-1436	.3032	. 2669	.2529	.2226	.2781	-2447
.273	.1442	.3043	. 2679	.2538	.2234	.2791	. 2456
.274	-1447	-3054	. 2688	-2547	.2242	.2801	. 2465
•275	.1452	.3065	. 2698	.2556	.2250	.2811	-2473
.276	-1457	.3076	-2707	.2565	.2258	.2821	.2482
-277	. 1462	.3086	.2717	-2574	. 2266	. 2830	.2490
.278	.1467	.3097	.2726	.2583	.2274	. 2840	.2499
-279	-1473	.3108	.2736	.2592	.2282	.2850	.2508
.280	.1478	.3119	.2745	. 2602	.2290	.2861	.2517
.281	.1483	.3130	-2755	.2611	.2298	.2871	. 2526
.282	.1488	.3140	.2764	. 2620	.2306	.2880	-2534
. 283	-1493	.3151	-2774	. 2629	.2314	. 2890	.2543
. 284	.1498	.3162	. 2783	. 2638	.2322	.2900	.2552
.285	.1504	-3173	.2793	.2647	.2330	.2910	.2561
. 286	.1509	.3184	. 2802	.2656	.2338	.2920	.2570
.287	.1514	.3194	. 281 2	.2665	.2346	.2930	.2578
. 288	.1519	.3205	.2821	.2674	.2354	. 2940	.2587
.289	.1524	.3216	. 2831	.2683	.2362	. 2950	.2596
.290	.1529	.3227	. 2840	. 2693	.2370	.2960	.2605
.291	-1535	.3238	. 2850	.2702	.2378	.2970	.2614
.292	.1540	.3248	.2859	. 2711	.2386	.2980	.2622
•293	-1545	-3259	. 2868	.2720	-2394	.2990	.2631
•294	.1550	-3270	. 2878	. 2729	.2402	.3000	.2640
.295	-1555	.3281	. 2887	.2738	.2410	.3010	.2649
. 296	.1560	.3292	. 2897	-2747	.2418	.3020	.2658
- 297	.1566	.3302	.2906	.2756	.2426	.3030	.2666
. 298	.1571	-3313	.2916	.2765	-2434	.3040	.2675
.299	.1576	-3324	-2925	-2774	.2442	.3050	. 2684
.300	.1581	-3335	-2935	.2784	.2450	.3060	.2693

SEPARATION AND DETERMINATION OF THE VARIOUS CARBOHYDRATES OF CEREALS, ETC. STONE'S METHOD.

Stone has thus tabulated the results of a series of analyses of various samples of wheat, flour, corn, and bread, in which he has separated the principal carbohydrates.*

PERCENTAGES OF VARIOUS CARBOHYDRATES IN CERTAIN FOODSTUFFS.

	Sucrose.	Invert Sugar.	Dextrin.	Soluble Starch.	Pento- sans.	Crude Fiber.
Whole wheat, I	0.52	0.08	0.27	0.00	4.54	2.68
Whole wheat, II		0.00	0.41	0.00	4-37	2.51
Wheat flour, I		. 0.00	0.00	0.00	0.00	0.25
Wheat flour, II		0.00	1.06	0.00	0.00	0.25
Corn	9.27	0.00	0.32	0.00	5.14	1.99
Sugar-beet		0.07	0.35	0.00	4.80	1.00
Bread (wheat, I)		0.32	0.68	1.37	4.16	2.70
Bread (wheat, II)		0.37	0.23	2.36	4.34	2.02
Bread (flour, I)		0.10	0.27	1.00	0.00	0.34
Bread (flour, II)		0.38	0.91	1.74	0.00	0.17
Corn cake (maize)		0.19	0.00	2.80	3 - 54	2.22
•	1	_	l	1		l

Determination of Cane Sugar.—100 grams of the finely ground material are extracted by boiling under a reflux condenser with 500 cc. of 95% alcohol for three hours, the alcoholic extract is filtered, evaporated nearly to dryness, and then taken up with a small amount of water, to separate the sugar from the oils and waxes dissolved by the alcohol. This aqueous solution is invariably dextro-rotary, and seldom contains any reducing sugar. If the latter is present, it is determined in an aliquot part of the aqueous solution with Fehling's solution, the result being calculated to dextrose. The remainder of the aqueous sugar solution, or the whole of it, if, as is almost always the case, dextrose is absent, is then inverted by heating with hydrochloric acid in the usual manner (page 588) and the sugar is estimated with Fehling's solution, calculating the result to sucrose (page 612).

Determination of Dextrin.—Digest the residue from the above alcoholic extraction from eighteen to twenty-four hours with 500 cc. of cold distilled water, shaking frequently. On filtering, a clear solution is ob-

^{*} Jour. Am. Chem. Soc., 19, 1897, p. 183, and U. S. Dept. of Agric., Off. of Exp. Sta., Bul. 34. The percentages of normal starch found by Stone are obviously erroneous, and are for this reason excluded from the table as here given.

tained, which should be tested with iodine for soluble starch. If the latter is not found (which is nearly always the case), the solution is concentrated to a small volume, avoiding a temperature higher than 80° to 90°, and this is boiled under a reflux condenser for two hours with one-tenth its volume of hydrochloric acid (specific gravity 1.125). Determine the dextrose by Fehling's solution and calculate to dextrin by the factor 0.9. Or, instead of submitting the concentrated aqueous extract to hydrolysis as above, the dextrin may be roughly determined gravimetrically therein by treating with several volumes of strong alcohol until no further precipitation is produced. The flocculent precipitate thus obtained is collected, dried, and weighed.

Determination of Starch.—Dry in an oven the residue from the preceding treatment and determine its quantitative relation to the original sample; 2 grams are then accurately weighed and subjected to the diastase method of starch determination (page 283).

Determination of Pentosans and Hemicelluloses.—The washed residue, left after filtering off the starch-containing solution from the process of heating with malt extract in the preceding starch determination, is boiled for an hour with 100 cc. of 1% hydrochloric acid, which converts all the pentosans into sugar. Filter, and wash the residue thoroughly, make up the solution to 200 cc., and determine the sugar with Fehling's solution, calculating the results for xylan, assuming that the chief sugar formed is xylose. The reducing power of xylose is assumed to be 4.61 milligrams for each cubic centimeter of Fehling's solution. If the volumetric Fehling method is used, 10 cc. of Fehling's solution are thus equivalent to 0.046 gram xylose. Xylose×0.88=xylan.

Crude Fiber (Cellulose, etc.).—The residue from the last dilute acid hydrolysis is boiled with 200 cc. of 1.25% solution of sodium hydroxide for half an hour, filtered, dried, and weighed. It is then ignited, and the weight of the ash deducted from the first weight.

PROTEINS OF CEREALS AND VEGETABLES.

Different cereal and vegetable foods present considerable variations in the character and extent of their protein constituents, and by no means all of the common vegetable foods have been studied in detail.

Osborne, in connection with Voorhees and Chittenden, has made a careful study of the proteins of many of the cereals, of potatoes, and of peas. A brief outline only will be given in what follows of methods

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for separation of the vegetable proteins. For fuller details the reader is referred to the work of Osborne et al. in the American Chemical Journal, Vols. 13, 14, and 15, and to the Journal of the American Chemical Society, Vols. 17, 18, 19, and 20.

Proteins Soluble in Water and Dilute Salt Solution.—By the action of various solvents it is possible to separate the different classes of proteins for examination or analysis. Thus water at first applied extracts certain of the soluble proteins, as does a weak salt solution. Osborne and Voorhees recommend the use of a 10% solution of sodium chloride as the first solvent to apply for separating vegetable proteins, shaking the finely ground material with twice its weight of the salt solution. The salt solution, after filtering, is then subjected to dialysis, the protein matter thus separated out being a globulin, while that not precipitated on dialysis is assumed as the protein matter of the substance soluble in water. Two albumins and a proteose are found in wheat to be thus soluble in water.

If the proteins soluble in salt solution are to have their total nitrogen determined, they are completely precipitated from the solution by saturating with zinc or ammonium sulphate.

There are thus two classes of proteins soluble in 10% salt solution: (a) globulins, insoluble in water alone, and (b) albumins and proteoses, which are soluble in water.

Separation of Albumins, Proteoses, and Globulins.—Starting with the aqueous solution containing the albumins and proteoses, if present, the former are best separated according to Osborne and Vorhees by fractional coagulation, effected by heating at different temperatures, those that precipitate out at a temperature under 65° being first filtered out, and the filtrate submitted to a higher temperature not exceeding 85°. The two portions thus separated may be collected in filters, and their nitrogen separately determined.

The proteose may be precipitated from the filtrate by saturating with ground salt, or by adding, first salt to the extent of 20%, and finally acetic acid.

The globulins, precipitated in the original 10% salt solution by the process of dialysis as described, may themselves be separated by employing salt solution of varying strength as solvents.*

Proteins Soluble in Dilute Alcohol, but Insoluble in Water.—The residue from the treatment with 10% sodium chloride is digested with 75% alcohol at about 46° C. for some time and filtered. The residue is further

^{*} Am. Chem. Jour., 13, p. 464.

digested at about 60° with 75% alcohol three separate times. The evaporated filtrates contain the alcohol-soluble proteins. In this class are the hordein of barley, the gliadin of wheat and rye, and the zein of corn.

Proteins Insoluble in Water, Salt Solution, and Dilute Alcohol.—It is customary to determine the nitrogen in the final residue without further attempt to separate the remaining protein matter. It is, however, possible to further extract with alkaline and acid solvents, if desired, which process, however, changes the nature of the proteins from that in which they originally exist in the substance.

Character and Amount of Proteins in Wheat.*—The proteins of wheat, according to Osborne, are five in number, as follows:

	A	mount Present, Per Cent.
Soluble in water:	Albumin (leucosin) Proteose	0.3 to 0.4
	Proteose	0.3
Soluble in 10 per cent NaCl:	Globulin (edestin)	0.6 to 0.7
Soluble in dilute alcohol:	Gliadin	4.25
Insoluble in above:	Glutenin	4.00 to 4.5

The term gluten is applied to the protein content of wheat flour insoluble in water, the value of flour for baking bread depending on the amount present. Gluten contains the two definite proteins, gliadin and glutenin. Crude gluten, as obtained by washing the dough in the analytical process (page 319), is a complex mixture of many bodies, containing, besides the two proteins above named, small quantities of cellulose, mineral matter, lecithin, and starch.

Separation and Determination of Wheat Proteins.—Teller's Method.†—Non-gluten Nitrogen.—Two grams of the finely divided sample are mixed with about 15 cc. of 1% salt solution in a 250-cc. flask. The flask is shaken at intervals of ten minutes during one hour, after which it is filled to the mark with the salt solution and allowed to stand two hours. The supernatant liquid is then filtered through a dry filter into a dry flask, leaving most of the solid material in the flask, passing the first part through twice, if necessary, for a clear filtrate. With a pipette, exactly 50 cc. of clear filtrate are run into a 500-cc. Kjeldahl digestion-flask, 20 cc. of the usual reagent sulphuric acid for the Gunning process (p. 69) are added, and the contents of the flask brought to a gentle boil. After the water has

^{*} Am. Chem. Jour. XV, 392-471; XVI, 524. †'Ark. Exp. Sta. Bul. 42, p. 96.

been driven off and the acid has stopped foaming, the potassium sulphate is added and the digestion completed. From the per cent of nitrogen thus obtained 0.27% is deducted, this figure corresponding to the amount of gliadin soluble in 1% salt solution under the above conditions. The remainder is the percentage of non-gluten nitrogen.

Gluten Nitrogen.—This is obtained by difference between the total nitrogen and the non-gluten nitrogen as above obtained, or by deducting the combined nitrogen of the edestin, leucosin, and the amido-nitrogen from the total nitrogen.

Edestin and Leucosin.—Edestin is a globulin belonging to the vegetable vitellins, and is precipitated from salt solutions by dilution, or by saturation with magnesium or ammonium sulphate, but not by saturating with sodium chloride. It is not coagulated below 100° C., but is partly precipitated by boiling. Leucosin is an albumin, coagulating at 52°, but precipitates from salt solution by saturating with sodium chloride or magnesium sulphate.

To 50 cc. of the clear salt extract, obtained as described under non-gluten nitrogen, 250 cc. of pure 94% alcohol are added in a Kjeldahl 500-cc. digestion-flask, the contents thoroughly mixed, and allowed to stand over night. The precipitate is collected in a 10-cm. filter, which is returned to the flask and the nitrogen determined. This represents the nitrogen of the combined edestin and leucosin. These proteins may, however, be separated by coagulating the leucosin at 60°, and precipitating the edestin by adding alcohol to 50 cc. of the clear filtrate, determining the nitrogen separately in each precipitate.

Amido-nitrogen.—Allantoin, asparagin, cholin, and betaine are nitrogenous bases present in wheat.

Ten cc. of a 10% solution of pure phosphotungstic acid are added to 100 cc. of the clear salt extract as above obtained, thus precipitating all the proteins, which are allowed to settle preferably over night. Filter, and determine the nitrogen in the clear filtrate. The filtrate should be tested with a little of the phosphotungstic acid reagent to make sure that all the proteins have been separated. In some cases, as in bran for instance, more than 10 cc. of the reagent are necessary.

Gliadin is dissolved most readily from flour by hot dilute alcohol, but is entirely insoluble in absolute alcohol. One gram of the material is extracted with 100 cc. of hot 75% alcohol, by shaking the mixture thoroughly in a flask, and heating for an hour at a temperature just below

the boiling-point of alcohol, with occasional shaking. After standing for an hour, the hot liquid is decanted upon a 10-cm. filter, and 25 cc. of the hot alcohol are added to the residue and shaken, after which the residue is again allowed to settle, and the liquid decanted. This is repeated six times. The remainder of the alcohol is then driven off by evaporation, and the nitrogen determined in the residue. The difference between the total nitrogen and the nitrogen thus obtained, gives the nitrogen of the alcoholic extract, which includes the amides. Subtracting the latter, or amido-nitrogen, the remainder is the gliadin nitrogen.

Glutenin Nitrogen.—This is the difference between the gluten nitrogen and the gliadin nitrogen.

The factor by which the nitrogen should be multiplied in determining the various proteins, according to Osborne and Voorhees, is 5.7 for wheat.

Proteins of the Common Cereals and Vegetables.—Osborne and his coworkers have made a detailed study of the protein constituents not only of wheat as above outlined, but of other common grains and vegetables, and the results of these investigations may be thus briefly summarized:

Proteins of rye: *	
Insoluble in salt solution	1.00
Proteins of barley:†	ent.
Soluble in water: { Leucosin } Proteose }	5.3
Soluble in salt solution, edestin	1.00
Proteins of corn:	
Soluble in water: Proteose	0.04
Protein of pea:§	-
Soluble in salt solution: Globulins { Legumin	7.00 3.00 2.03

^{*} Jour. Am. Chem. Soc., 17, page 429. † Ibid., 17, p, 539.

Proteins of Potato.*—Almost the whole protein content of the potato consists of a globulin to which Osborne has applied the name "tuberin." Proteose is also present in very small amount.

MINERAL CONSTITUENTS OF CEREALS AND VEGETABLES.

The food analyst often finds the determination of one or more of the mineral constituents of a food product of value as a means of detecting adulteration, since the addition of foreign material may alter materially the composition of the ash.

The table† on page 302 shows the composition of the pure ash of common cereals.

Scheme for Complete Ash Analysis.—The following scheme in essential details was suggested by the late Prof. S. L. Penfield of Yale University, and has been in use for over twenty years at the Connecticut Agricultural Experiment Station.

Preparation of Ash.—The amount of material which should be reduced to ash depends on the percentage of total ash present and the amount of material available. Usually 100 grams is a suitable amount; if, however, the material (e.g., tobacco) is rich in ash, 50 grams is sufficient, while if it contains but a small amount of ash, 200 grams or even more may be required. About 5 grams of ash is a liberal amount for a complete analysis, but in case of necessity 1 gram will suffice if care is taken to so adapt the scheme as to make as many determinations as possible on one weighed portion.

The ashing is carried on in a platinum dish heated below redness by a Bunsen burner. In order to distribute the heat and prevent overheating, a piece of asbestos paper is introduced between the dish and the flame. The material first chars, then begins to glow just below the surface, and the combustion gradually extends downward until it reaches the bottom of the dish. Then, and not until then, the unburned carbon on the surface should be stirred in with the ash to facilitate burning. Care should be taken not to heat higher than dull redness, thus avoiding the loss of alkali chlorides and the fusion of alkali phosphates about the particles of carbon. A muffle furnace may be used to complete the burning.

Substances rich in starch or sugar are most difficult of combustion,

^{*} Jour. Am. Chem. Soc., 18, 1896, p. 575.

[†] U. S. Dept. of Agric., Bur. of Chem., Bul. 13, part 9, p. 1212.

COMPOSITION OF ASH OF CEREALS.

	K.O.	Na ₂ O.	CaO.	MgÖ.	Fe ₂ O ₂ .	PeOg.	SO ₃ .	C1.	SiO ₂ .
Wheat (Canada) Rye (Minnesota). Barley (U. S.) Oats (U. S.) Corn (U. S.). Rice, polished (Guatemala). Buckwheat (U. S.)	27.60 24.15 15.91 33.92 20.84	4.64 6.42 4.38 7.72 13.98	5.56 2.44 4.09 3.18 4.48	11.73 8.23 7.18 17.99	5.23 0.33 0.20 0.50 0.89	35·47 24·34 35·25 43·21	0.52 0.22 0.48 0.44	0.58 0.56 1.02 0.00 0.80	2.45 22.30 42.64 1.00 6.14

Teller * obtained the following results of ash analyses of flour, bran, and wheat:

ASH OF WHEAT PRODUCTS.

	Patent Flour.	Straight Flour.	Low Grade.	Bran.	Wheat,
Silica	2.33	1.28	.50	•97	1.04
Alumina	.41	.15	.12	.07	.11
Ferric oxide	-47	.26	.25	-27	.27
Potash	38.50	36.31	32.27	28.19	29.70
Soda	0.00	0.00	0.00	0.00	0.00
Lime	5-59	5.65	4.51	2.50	3.10
Magnesia	4-39	6.44	9-33	14.76	13.23
Phosphoric acid	48.05	49-32	53.10	52.18	52.14
Sulphur trioxide	.16	-52	.00	.10	.22
Chlorine	• • • • • •		•••••	.01	.01
Zinc oxide	•••••	.04	•••••	-27	-24
Sum	99.90	99-97	100.08	99-95	100.06
Per cent of total ash	-31	.40	.70	•••••	1.62

König gives the following analyses of the ash of various leguminous and other vegetables:

	Number of Analyses.	Ash in Dry Substance.	Potash.	Sode.	Lime.	Magnesia.	Iron Oxide.	Phosphoric Acid.	Sulphuric Acid.	Silica.	Chlorine.
Beans	15	3.57	42-49	1.34	4-73	7.08	o.57 o.86	38.74	2.53	0.73	1.57
Peas Potatoes	53	3-77	41.79	0.96 2.62	4-99		1.18			0.86 2.13	3.11
Beets	15	6.44	54.02		4.12				3.17	2.38	8.40
Carrots	11	5-58	35.21	22.07		4-73	1.03	12.46	6.72		
Turnips	32	8.01	45-40	9.84	10.60	3.69	0.81	12.71	11.19	1.87	5.01

^{*} Ark. Exp. Sta. Bul. 42.

as the charcoal forms a hard mass, while substances rich in fibrous or woody matter burn quite readily without losing their powdered condition. A certain amount of unburned carbon is no disadvantage, as it is determined in the course of the analysis.

Finally cool the ash, grind to a powder, mix without loss, and weigh, thus determining the percentage of crude ash.

Determination of Water.—Heat I gram of the ash in a platinum crucible well below redness to constant weight.

Determination of Carbonic Acid. — Determine carbonic acid as described on p. 336 using the portion dried for the determination of water.

Determination of Charcoal and Sand.—Weigh I gram of the ash, or transfer the solution and residue from the determination of carbonic acid, into a beaker, add 25 cc. of water and 25 cc. of 10 per cent hydrochloric acid, and boil gently for 10 minutes. Filter on a Gooch crucible, and wash thoroughly with hot water. Reserve the filtrate for determination of silica, iron oxide, alumina, lime, and magnesia. Wash the residue on the crucible once with alcohol and once with ether, and dry to constant weight at 100° C. Ignite and weigh again. The loss on ignition is the charcoal, the residue is sand.

Determination of Silica, Iron Oxide, Alumina, Lime and Magnesia.— Evaporate to dryness in a platinum dish the filtrate from the determination of charcoal and sand, heat for some hours on the water bath, and dry at 130° C. until all hydrochloric acid is removed. Moisten the residue thoroughly with concentrated hydrochloric acid, add hot water, stir, and decant the solution on an ashless filter. Treat the residue again with acid and hot water, and repeat the treatment until nothing but silica remains undissolved. Finally collect the silica on the paper, wash with hot water, ignite in a platinum crucible, and weigh.

To the filtrate add ammonia until a precipitate forms which remains on stirring, and then add sufficient hydrochloric acid to just dissolve the precipitate. Heat to 50° C. and add an excess of ammonium acetate solution and 4 cc. of 80 per cent acetic acid. Digest at 50° C. until the mixed phosphates of iron and alumina have settled, filter, wash with hot water, ignite in a platinum crucible, and weigh. As the precipitate is usually slight and consists almost entirely of iron phosphate, the iron oxide may be calculated with reasonable accuracy using the factor 0.53. If, however, greater accuracy is desired fuse the weighed precipitate with 10 parts of sodium carbonate, dissolve in dilute sul-

phuric acid, reduce with hydrogen sulphide, determine iron by the volumetric permanganate method, and in the same solution determine phosphoric acid by the molybdic method. The alumina is obtained by difference, subtracting the sum of the weights of the oxide of iron and phosphoric acid from the total weight of the precipitate.

To the filtrate from the mixed phosphates add an excess of ammonium oxalate, allow to stand in a warm place over night, filter, ignite the precipitate in a platinum crucible over a Bunsen burner, and finally to constant weight over a blast lamp, thus obtaining the calcium oxide.

Precipitate the magnesia in the filtrate from the lime by adding ammonia to alkaline reaction, then an excess of sodium phosphate solution with constant stirring, and finally sufficient concentrated ammonia to form one-tenth the final volume. Let stand over night, collect the magnesium ammonium phosphate on a Gooch crucible, ignite to magnesium pyrophosphate, and weigh.

Determination of Sulphuric Acid, Potash, and Soda.—Boil I gram of the ash with dilute hydrochloric acid, and remove charcoal, sand, and silica, as described in the preceding section. Evaporate nearly to dryness to remove the excess of acid. Dilute to 100 cc., heat to boiling, and add barium chloride solution drop by drop until the sulphuric acid is precipitated. Allow to stand over night, filter, ignite, and weigh as BaSO₄.

Heat the filtrate to boiling, add enough barium hydroxide to make the solution strongly alkaline, filter, and proceed with the determination of potash and soda, as described on p. 345.

Determination of Phosphoric Acid.—Dissolve 0.5 gram of the ash in hydrochloric acid, filter, and wash. Neutralize with ammonia, clear with nitric acid, and proceed as described on p. 346.

Determination of Chlorine.—Dissolve I gram of the ash in cold, very dilute nitric acid, filter, and wash. To the filtrate add an excess of silver nitrate, and heat nearly to boiling with constant stirring. Filter on a Gooch crucible, wash with hot water, dry the precipitate at a low heat, and heat cautiously at dull redness until the silver chloride has partially melted.

If desired the chlorine may be determined volumetrically by Volhardt's method, as follows: To the nitric acid solution add a known volume of decinormal silver nitrate solution sufficient to precipitate the chlorine, and 5 cc. of saturated solution of ferric alum. Titrate with decinormal ammonium thiocyanate solution until a permanent brown

color is formed. Subtract the volume required from the volume of decinormal silver nitrate added, and calculate the chlorine.

Determination of Sulphur in Vegetable Materials.*—Place from 1.5 to 2.5 grams of material in a nickel crucible of about 100 cc. capacity. and moisten with approximately 2 cc. of water. Mix thoroughly, using a nickel or platinum rod. Add 5 grams of pure anhydrous sodium carbonate, and mix. Add pure sodium peroxide, small amounts (approxmately 0.50 gram) at a time, thoroughly mixing the charge after each addition. Continue adding the peroxide until the mixture becomes nearly dry and quite granular, requiring usually about 5 grams of peroxide. Place the crucible over a low alcohol flame (or other flame free from sulphur), and carefully heat with occasional stirring until contents are fused. (Should the material ignite the determination is worthless.) After fusion, remove the crucible, allow to cool somewhat, and cover the hardened mass with peroxide to a depth of about 0.5 cm. Heat gradually, and finally with full flame until complete fusion takes place. rotating the crucible from time to time in order to bring any particles adhering to the sides into contact with the oxidizing material. Allow to remain over the lamp for ten minutes after fusion is complete. Cool somewhat. Place warm crucible and contents in a 600 cc. beaker. and carefully add about 100 cc. of water. After violent action has ceased. wash material out of crucible, make slightly acid with hydrochloric acid (adding small portions at a time), transfer to a 500 cc. flask, cool, and make to volume. Filter, and take a 200 cc. aliquot for determination of sulphates by precipitating with barium chloride in the usual manner.

Determination of Chlorine in Vegetable Substances.* — Impregnate 5 grams of substance in a platinum dish with 20 cc. of a 5 per cent solution of sodium carbonate, evaporate to dryness, and ignite as thoroughly as possible. Extract the residue with hot water, filter, and wash. Return to the platinum dish, ignite to an ash, dissolve in nitric acid, and determine chlorine by the Volhard method (p. 304).

MICROSCOPY OF CEREAL PRODUCTS.

A study of the histology of the various cereal grains is beyond the scope of the present work, and the reader who wishes to pursue this branch of the subject is referred to the special treatises on this subject.

^{*}A. O. A. C. Method, U. S. Dept. of Agric., Bur. of Chem., Bul. 107 (rev.), pp. 23, 24. † Ibid, p. 24.

The characteristics of the tissues of the various ground cereal grains are quite distinctive, when carefully studied, sufficiently so, at least, to serve to identify the particular grain from which a given product is made. In the case of flour and some other products, however, the tissues are largely removed in the process of milling, and such fragments as remain are so small as to render identification difficult. Products of this nature are identified either by the character of the starch grains—as for example, in the detection of wheat or corn flour in buckwheat flour—or else, if the starch is not sufficiently characteristic—as in the detection of wheat flour in rye flour—by examining the tissues accumulated from a considerable amount of the material after removal of the starch.

We have already seen, that the various cereal starches differ considerably in morphology and mode of grouping from each other, and this is true to such an extent that the expert can readily identify them. Since starch furnishes much more than half the content of all ground cereals, any considerable admixture of one flour with another is nearly always rendered apparent by a careful study of the magnified starch grains, which form a large part of the field when viewed under the microscope.

The most convenient method of accumulating the tissues from flour is to mix thoroughly 2 grams of the material with 200 cc. of water and 2.5 cc. of sulphuric acid, bring to a boil, allow to settle, and carefully decant off the liquid. After transferring to a capsule by means of a little water, and removal of the latter by decantation, the tissues are mounted for examination in very dilute potassium hydroxide solution.

Wheat Products.—Fig. 62 and Pl. VIII show the principal elements of the wheat kernel.

The outer layer or epicarp (Fig. 62, epi¹ and epi²) consists of beaded cells, which on the body of the kernel are elongated, but at the end are polygonal. From this layer at the end of the kernel arise the hairs (Fig. 62, t, Pl. VIII, Fig. 151) which form a beard clearly visible under a lens. Some of these hairs become detached in milling, and pass endwise through the bolts, hence their presence in even the highest grade of flour. The second layer or hypoderm (hy) resembles the first, while the third, although likewise made up of beaded cells, is strikingly different and forms the most characteristic tissue of the grain. These cells (Fig. 62, tr, Pl. VIII, Fig. 150) being transversely extended are known as "cross cells," and are further distinguished from the outer layers by their arrangement side by side in rows. The cells of the intermediate layer

(Fig. 62, in) and the tube cells (tu^1 and tu^2), although of striking appearance, are not of as frequent occurrence as the other layers. The crossing layers of the seed coat or spermoderm (i and o), are often met with, and are characterized by the thin walls of the cells and their brownish color.

The perisperm (P), consisting of colorless cells, is seldom seen, except after special preparation, while the next layer, made up of aleurone cells

Fig. 62.—Wheat. Elements in Surface View. X160. (WINTON.)

epi¹ epicarp at end of grain, with t hairs; epi² epicarp on body of grain; hy hypoderm (first layer of mesocarp); in intermediate cells; tr cross cells; tu¹ typical tube cells; tu² tube cells passing into spongy parenchyma; o outer layer of spermoderm; i inner layer of spermoderm; P perisperm; of aleurone cells; om starch grains.

(Fig. 62, al; Pl. VIII, Fig. 150), is the most conspicuous of the kernel. This layer is not, however, characteristic of wheat, as it is found in all cereal grains and in buckwheat. The aleurone cells do not contain, as was formerly supposed, the gluten of the grain; this occurs with the starch in the thin-walled cells within the aleurone layer.

The starch granules (Fig. 62, am; Pl. VIII, Fig. 152) are described

on page 281. The starch cells and the aleurone cells together form the endosperm.

The germ, situated at one side of the lower end of the kernel, is made up of very small cells containing fat and protein, but no starch.

Rye Products.—The structure of rye (Fig. 63; Pl. VII) resembles closely that of wheat. The number and general characters of the cell layers are the same in both, and the starch granules are very much alike. There are, however, certain points of difference which serve to dis-

Fig. 63.—Rye, Outer Bran Layers in Surface View. Epicarp consists of porous cells with the hairs, and their scars; quecross cells. × 160. (MOELLER)

tinguish the products of the two cereals, and even to detect the presence of a wheat product in a rye product, and vice versa:

First. The breadth of the cavities of wheat hairs is usually less than the thickness of the walls, whereas in rye hairs the reverse is often true (Figs. 62 and 63, t).

Second. The cross cells of wheat have rather thick, distinctly beaded side walls, and thin, pointed end walls; the cross cells of rye have rather thin, indistinctly beaded side walls, and usually swollen, rounded end walls (Figs. 62 and 63, tr; Figs. 150 and 146).

Third. The large starch granules of wheat seldom reach 0.050 mm. in diameter, while those of rye frequently exceed that limit. Radiating clefts often occur in the starch granules of rye (Pl. VII, Fig. 148).

Fourth. Wheat flour yields a considerable amount of gluten when

treated according to Bamihl's test (page 322); rye flour yields none or only a trace.

Barley Products.—The common varieties of barley are "chaffy," that is, the grain after threshing is still closely invested by the chaff (Pl. I, Fig. 123). The grain within the chaff is analogous in structure to wheat and rye, but differs from these in that the cross cells are not beaded and form a double layer (Fig. 64, tr), and the starch granules

Fig. 64.—Barley. Surface view of tr double layer of cross-cell; tw tube cells; is seed coat.

× 300. (MOELLER.)

seldom exceed 0.035 mm. in diameter (Pl. I, Fig. 124). The starch is more fully described on page 281.

Corn Products.—The most characteristic element of corn is the starch (page 281; Pl. IV). Polygonal starch granules 0.017 to 0.030 mm. in diameter occur in no other vegetable product of economic importance, excepting the seeds of Kaffir corn and other grains belonging to the genus Sorghum, which are used chiefly for cattle or poultry foods.

Oat Products.—The oat kernel resembles barley in appearance, but is not ribbed. In the preparation of oat meal and other breakfast foods, the chaff (Pl. IV, Fig. 135; Pl. V, Fig. 137) is removed and utilized as a cattle food. The elements of the grain of chief value in identification are the hairs and the starch granules. The hairs (Pl. V, Fig. 138) are much longer than those of wheat, rye, and barley, often reaching 1 mm. They taper toward both ends, so that when detached they often appear to be pointed at the base as well as at the apex. The

starch granules are small, of the polygonal type, and often occur in egg-shaped aggregates (page 282; Pl. V, Fig. 139).

Rice Products.—The chaff which envelopes this grain is rough and silicious, and after removal from the inner kernel is not suited even for

epi fu

es

N

Fig. 65.—Rice. Bran coats in surface view *epi* epicarp; *mes* mesocarp; *tr* cross cells; the tube cells, S seed coat; N perisperm. \times 300. (Winton.)

cattle food. Its appearance under the microscope is shown in Plate VI, Fig. 142. The thin skin of the kernel proper is largely but not entirely

Fig. 66.—Buckwheat. Bran coats in surface view. Seed coat consists of o outer epidermis, m spongy parenchyma, and ep inner epidermis; al aleurone cell. ×300. (MOELLER.)

removed in the preparation of rice for the market. The elements of this skin are shown in Fig. 65, the outer layer (epi) being the most characteristic. Unlike wheat, rye, and barley, it has no beard. Rice starch (Pl. VI, Fig. 143) is hardly distinguishable from oat starch. It is described on page 282.

Buckwheat Products.—In the preparation of buckwheat flour the black outer hulls and the inner skin or bran are largely, but not completely, removed. The bran elements are characteristic constituents of the flour, and are rendered especially distinct by adding a drop of dilute potassium hydroxide solution to a water mount (Fig. 66). The cells with wavy walls (0) and the spongy parenchyma (m) are especially noticeable. The starch of buckwheat resembles that of oats, but the individual granules are somewhat larger and occur in rod-shaped, not egg-shaped, aggregates (page 281; Pl. II, Fig. 128). Masses of starch granules (Pl. III, Fig. 129) conforming to the shape of the cells, occur in abundance in the flour.

RI.OIIR.

Flour is the term applied to the finely ground and bolted substance of wheat and other grains, though, unless otherwise qualified, by the term "flour" is generally understood that of wheat. In the process of manufacture, the dried wheat or grain is first crushed between mill-stones, forming the comparatively coarse product known as whole meal. This, by bolting, may be separated simply into flour and bran, but in the crude milling of years ago at least three products were usually obtained from wheat, viz., fine flour, coarser shorts, or middlings, and bran.

In the improved modern processes of milling, which meet the demands for the very finest flour, as well as other grades, the material is subjected to repeated sifting and grinding between grooved rollers, with the result that it is possible to turn out as many as ten separate grades of flour, as is shown by the following record of a mill near Trieste:

Groats.* A and B	2 per cent)
Flour, No. o	5 "	1
" " I	12 "	
" " 2	6 "	41 per cent extra flour
" " 3	6 "	42 For some came 2000
" " <u>4</u>	. «	
" " "	2 "	<u> </u>
" " 6	14 "	í
« « ₇	- "	
« « ģ	9 "	38 per cent medium and common
" " 0	J "	1
Bran	18 "	1
Loss	3 "	21 per cent waste
ANGS) =
	100 per cent	
	TOO DOT CETT	

* Masses of the interior of the berry.

	Analyses	of	these	separated	products	made	by	C.	A.	Pillsbury	are
as	follows:									•	

	Water.	Ash.	Phosphoric Acid.	Nitrogen.	Proteins Calculated
Groats	10.57	0.42	0.20	2.24	14.65
No. o	10.37	0.43	0.14	1.68	10.76
" I	10.23	0.41	0.21	r.68	10.76
" 2	10.47	1.03	0.22	1.72	11.02
" 3	10.07	1.02	0.17	1.72	11.02
" 4	70.24	1.19	0.25	1.74	11.15
" 5	9.66	0.69	0.35	1.80	11.54
" ő	11.12	1.04	0.24	1.84	11.79
7	10.99	0.81	0.21	1.80	11.54
* 8	9.86	1.01	0.36	1.90	12.18
9, coarse bran	9.71	7 - 32	2.14	1.98	12.60
" 10, fine bran	11.01	4.21	0.70	2.20	14.16

In this country the common practice of most mills is to produce about three grades of flour, somewhat as follows:

Graham Flour, or whole-wheat flour, is made from the unbolted meal of wheat, ground as finely as possible. It is actually a mixture of flour and bran.

Composition of Common Flours.—The following analyses are collated and summarized from Bulletin 13, part 9 of the Bureau of Chemistry:

	No. of Analy- ses.	Moisture.	Proteins, N×6.25.	Proteins, N×5.70.	Moist Gluten.	Dry Gluten.
Patent wheat flour	40	12.77	10.55	9.62	25.97	9.99
Common market wheat flour		12.28	10.18	9.28	24.55	9.21
Bakers' and family flour	14	11.69	12.28	11.20	34.70	13.07
Indian-corn flour	3	12.57	7.13	i		
Rye flour	I	11.41	13.56			
Barley flour	1	10.92	7.50			
Buckwheat flour		11.89	7.50 8.75			

	Ether Extract (Fat).	Ash.	Nitrogen- free Extract (Starch, Sugars. Gums.etc.)	Crude Fiber.	Calculated Calories of Combus- tion.
Patent wheat flour. Common market wheat flour. Bakers' and family flour. Indian-corn flour. Rye flour Barley flour Buckwheat flour		0.44 0.61 0.57 0.61 1.55 0.86 1.85	74.76 75.63 73.87 78.36 73-37 80.50 75.41	0.21 0.28 0.22 0.87 1.86 0.67 0.52	3858 3882 3929 3837

Damaged Flour.—Grain is often damaged by the growth of smuts, rusts and ergot. Both grain and flour are also liable to attacks of molds, yeasts, algae and bacteria.

Various insects and other forms of animal life frequently infest either the grain or the flour, or both. Among these are weevils and various other beetles, flour moths, mites, and the wheat worm, a nematode related to trichina.

Grain may also be damaged by sprouting, the diastase thus formed partially dissolving the starch granules with the formation of fissures and branching channels, which may be readily seen under the microscope. The amount of cold-water extract (page 320) shows to some extent whether or not a flour has been damaged.

Ergot.—Ergot is a fungus growth that occasionally develops within the grain of rye and other cereals, and, unless care is taken, becomes ground with them in the preparation of meal and flour. Ergot contains alkaloids of a poisonous nature, and instances are on record of its accidental presence in cereal preparations causing serious injury to health. While most commonly found in rye, ergot occasionally grows in wheat. If flour or meal containing ergot be treated with a very dilute solution of anilin violet, the stain will be practically absorbed by the damaged particles of the grain, and resisted by the normal granules. If shaken with dilute alkali, a meal or flour contaminated with ergot is colored violet, which by treatment with acid turns red. A hot, alcoholic extract of flour containing ergot is colored red when treated with dilute sulphuric acid.

Ergot in ground cereal preparations is best recognized under the microscope, appearing as a fine network of mostly colorless parenchyma cells, containing globules of fat (Fig. 67). Some of the cells are circular, others considerably elongated, and some contain a deep-brown coloring

matter, which, when treated with ammonia, takes a violet-red color. Occasionally the cell walls appear of a dark color. If the suspected sample be treated with dilute anilin violet, as above described, the stained ergot fragments will be especially apparent under the microscope.



Fig. 67.—A, Transverse Section of the Ergot of Wheat under the Microscope; B Powdered Wheat Ergot. (After Villiers and Collin.)

A

Adulteration of Flour.—Besides the substitution of cheaper or inferior grades for those of higher quality, the fraudulent admixture of cereals, other than wheat, is not uncommon in flour. Corn meal is sometimes found as an adulterant of wheat flour. Its presence is best detected by the microscope, the difference between the wheat and corn starch being readily apparent.

Rye flour is often adulterated with the cheaper grades of wheat flour or wheat middlings. This form of adulteration is detected by the Bamihl test (page 322) and by microscopic examination of the residue obtained after boiling with dilute acid (page 306).

Much of the so-called buckwheat flour consists of mixtures containing wheat or corn flour, or both. Rice flour is also used in pancake flours, although probably to not cheapen the product. Self-raising pancake flours are usually mixtures of two or more flours with leavening material.

The microscopic characteristics of the starch grains and tissues, serve to identify the different flours present in such mixtures.

Finely ground mineral adulterants are said to have been used in flours, but no authentic instance of this kind has come to the writer's knowledge. Any considerable admixture of such a nature would be manifest in the increased ash.

Alum in Flour.—The addition of alum to flour was formerly a common practice in Europe, both by miller and baker, for the purpose of improving the appearance of inferior or slightly damaged flour. Hence it was frequently found in the cheaper grades of flour and bread. Now, however, it is rarely if ever used for this purpose, and the presence of notable quantities of alum or its compounds in flour or bread is usually due to its use as an ingredient of leavening powders.

Detection.—To detect alum in flour, mix about 10 grams of the sample in 10 cc. of water and add about 1 cc. of an alcoholic tincture of logwood (5 grams logwood digested in 100 cc. alcohol) and about 1 cc. of a saturated solution of ammonium carbonate. Stir the whole well together. If the sample is pure, the color will be a faint brown or pink, but if alum is present, a distinct lavender-blue color is produced, which should remain after heating for two hours in the water-oven.

Alum may also be tested by the ammonium chloride method, described on page 344.

Bleaching of Flour.—Within the past few years various processes for bleaching flour with nitrogen peroxide have come into extensive use both in Europe and America. The nitrogen peroxide is generated by electrical, chemical, or electro-chemical means, and is diluted with air before treatment of the flour. In the Alsop process, which is most commonly employed, the gas is formed by a flaming discharge of electricity, which causes the nitrogen and oxygen of the air to combine.

Nitrogen peroxide destroys almost immediately the yellow color which is associated with the fat of the flour, thus increasing the whiteness of the product. It also combines with the moisture of the flour, forming nitrous and nitric acids, the nitrous acid (free or combined) being especially noteworthy because of the ease of detection. Snyder, after conducting baking tests, reached the conclusion that the bread made from bleached flour does not contain nitrite-reacting substances; Ladd, Mitchell, and Winton, however, have found that when made by the usual methods it contains appreciable quantities, and certain results obtained by Alway bear out this conclusion. Bleaching also diminishes the iodine number

of the fat of the flour, and is believed by some to affect the quality of the gluten.

Experiments of Shepard and of Ladd indicate that bleaching introduces toxic substances into the flour, although this is disputed by certain advocates of bleaching. Mitchell and also Winton find that bleaching injures the flavor of the bread.

Secretary Wilson of the U. S. Department of Agriculture issued a decision on December 9, 1908, declaring the bleaching of flour with nitrogen peroxide to be illegal under the Food and Drugs Act of June 30, 1906.

INSPECTION AND ANALYSIS OF FLOUR.

In some of the larger cities, authorized flour inspectors are appointed, whose business it is to examine the product and pass upon its quality. To such inspectors the local dealers submit samples, which the inspectors gauge as to color, soundness, weight, etc., comparing them usually with a series of graded samples, and stamping or branding them officially with the date as well as the grade. Market quotations also are based on the standard terms adopted. The names of the various grades differ with the locality. In St. Louis, the following names are adopted in order of their quality, viz., Patent, Extra Fancy, Fancy, Choice, and Family.

Such systems of inspection are under the auspices of local dealers, being organized and maintained for their own protection, and have not as yet been the subject of state or even municipal supervision, as in the case of meat inspection. The grade or quality of flour is determined largely by its appearance and color, by its fineness as indicated by rubbing between the fingers, by its odor, and by the so-called "doughing test," which consists in kneading the flour with water under fixed conditions, and noting its tenacity and elasticity. Baking tests are also relied on to a considerable extent by millers and buyers.

Of the chemical methods employed in grading flour, those for the determination of ash, protein, and gluten are of chief importance.

Fineness. — This is determined by rubbing the flour between the thumb and fingers. A gritty flour is one that is so coarsely ground that it feels rough and granular. When treated with water on a glass plate, the individual granules are evident. Examined under the microscope, the coarse granules are seen to consist of aggregates of cells, the contents of which are still intact. Smooth flour, on the other hand, feels soft and slippery. It is so finely ground that the cells are isolated and often

ruptured, a considerable part of the powder consisting of the starch grains and other cell contents, which have been liberated from the cells.

Pekar's Color-test.—Place 10-15 grams of the flour on a rectangular glass plate, about 12 cm. long and 8 cm. wide, and pack on one side in a straight line by means of a flour spatula. Treat the same amount of the standard flour used for comparison in the same manner, so that the straight edges of the two flours are adjacent. Carefully move one of the portions so as to be in contact with the other, and "slick" both with one stroke of the spatula, in such a manner that the thickness of the layer diminishes from about 0.5 cm. on the middle of the plate to a thin film at the edge, and the line of demarcation between the two flours is distinct. Cut off the edges of the layer with the spatula, so as to form a rectangle, and compare the color of the two flours. The difference in color becomes more apparent after carefully immersing the plate with the flour in water, and still more apparent after drying.

Absorption and Dough Test.—Stir 30 grams of the flour in a heavy coffee cup with 15 cc. of water by means of a spatula until a smooth ball of dough has been formed. If after standing two minutes the amount of water proves insufficient to thoroughly dough up the flour, repeat the operation, using 15.5 cc. of water, and, if necessary, continue to repeat until the quantity is found that will yield a stiff, but thoroughly elastic dough. From the results of this test, calculate the absorption of 1000 grams of flour in terms of cc. of water.

The physical characters of the dough, such as color and elasticity, furnish to the expert valuable indications of the quality or grade of the flour.

Expansion of Dough.—Rub to a smooth paste 3.5 grams of granulated sugar, 1.2 grams of salt, and 3 grams of compressed yeast, and thoroughly mix with 60 cc. of water at 35° C. Warm 100 grams of the flour in a shallow pan to 35° C, add to it the yeast mixture, mix with a spatula, and knead with the fingers until a smooth ball of dough has been formed. Drop the dough into a graduated, 500 cc. cylinder, pat down so as to force out the air, and note the volume of the mass. Place in a raising closet kept at 35° C. Read the volume at the end of the first hour and every half hour thereafter until the maximum is reached.

Baking Tests.*—Koelner's Method.—This process, also known as the straight dough method, yields a close-grained loaf of even texture, and serves well to determine the flavor and relative size of the loaf.

^{*} Descriptions by Miss H. L. Wessling, Chicago Laboratory, Bur. of Chemistry.

Place 220 grams of the flour, previously warmed in a shallow pan, in a raising closet kept at 35° C., in a Koelner dough kneader, which has previously been warmed to 35° C. by means of water placed in the special compartment for this purpose. To the flour add 12 grams of sugar, 5 grams of salt, and 10 grams of compressed yeast, rubbed smooth and thoroughly mixed in a cup with 100 cc. of water at 35° C. Rinse the cup with sufficient water to make the total quantity required, as calculated from the absorption test. This amount is usually about 87 cc.

Adjust the blades of the kneader for mixing, and turn the crank at the rate of 90 revolutions per minute for 10 minutes. Adjust the blades for kneading, add 120 grams of flour, previously warmed to 35° C., and turn the crank for ten minutes at the rate of 60 revolutions per minute. Remove the dough immediately to a warmed plate, cut into two equal parts, mould the two separately, and place end to end in a warmed, greased, and tared baking tin measuring 27×6.3 cm. at the top, 25.4×5 cm. at the bottom, and 8.8 cm. deep—all inside measurements.

Weigh the tin with dough, place a tin gauge across the top, and set the whole in the raising closet. After the dough has risen to the gauge, place the tin in a suitable oven heated to 200° C., and bake at 200 to 205° until 30 grams of water have been removed, which usually requires from 30 to 35 minutes. Break the loaf in two, and note the odor when hot and again when cold.

When thoroughly cool, determine the volume of the loaf as follows: cover the bottom of a box $7.6 \times 12.7 \times 28$ cm., inside measurements, with flaxseed, place the loaf in the box, pour flaxseed without jarring into the box until filled, and strike off the surplus seed by means of a straight edge. Remove the seed from the box, weigh, and divide the weight by the weight of I cc. of the seed, as calculated from an actual weighing of the seed required to fill the box. Subtract this figure from the cubic contents of the box in cc., thus obtaining the volume of the loaf.

Long Fermentation Method.—This method, used in some of the large mills in the northwest, differs from the Koelner method in that (1) a sponge is set, (2) the dough is kneaded twice, and (3) the dough is finally allowed to rise until it will allow no further expansion. The bread is coarse in texture, but serves well the purpose of determining the strength and flavor.

To 255 grams of warmed flour contained in a jar or earthenware crock, add 3.5 grams of salt and 8.6 grams of compressed yeast, mixed thoroughly with 170 cc. of water at 35° C. Stir together until a soft

sponge is formed, set in the raising closet, warmed to 35° C., and allow to rise until the volume has been doubled. Mix the risen sponge with 85 grams of warmed flour, 12 grams of sugar, 6 grams of lard, and the remainder of water (the total quantity for 340 grams of flour being calculated from the absorption test). Knead steadily for 6 minutes, transfer the dough to the jar or crock, and set it in the raising closet until it has again doubled its volume.

Remove the dough to a warmed plate, knead lightly in the hands for a minute or two, then place in a warmed, greased, and tared standard baking tin, which measures 16.8×8.8 cm. across the top, 14.9×6.9 cm. across the bottom, and 13.9 cm. deep, all inside measurements, and has wings or extensions of the metal at the top of the two sides. Weigh the pan and dough, prick the latter about a dozen times with a fine pointed wire, and set again in the raising closet. Let it rise in the pan until the maximum volume has been reached, *i.e.*, until the bubbles of gas just begin to break and form larger ones. This is a matter of judgment and can be learned only by experience.

Since the dough always rises somewhat in the oven, it must not be raised to its limit beforehand, but must be put in the oven at such a stage that with the additional rising in the oven it will have attained the maximum volume. A comparison of the volumes of different loaves will then be a means of judging the strength of the different flours. The loaf should be baked from 30 to 35 minutes, raising the temperature of the oven gradually from 180° C. at the beginning to 210° C. at the end.

Determination of Proximate Constituents of Flour. — Moisture, ash, protein, crude fiber, and fat are determined by the usual methods (pp. 277 to 279).

Determination of Moist and Dry Gluten.*—Place 25 grams of the flour in a coffee cup, add 15 cc. of water at a temperature not to exceed 15°, and work the mass into a ball with a spatula, taking care that none of it adheres to the dish. Allow the dough to stand one hour, then knead in a stream of cold water over a piece of bolting cloth held in place by two embroidery hoops, until the starch and soluble matters are removed. Place the gluten thus obtained in cold water, and allow to remain for one hour, after which press as dry as possible between the hands, roll into a ball, place in a tared flat-bottomed dish, and weigh as moist

^{*} U. S. Dept. of Agric., Bur. of Chem., Bul. 81, p. 118.

gluten. Spread the gluten out in the dish, dry for 24 hours in a boiling water oven, and weigh again, thus obtaining the dry gluten.

Determination of Alcohol-soluble Protein (Crude Gliadin).—Chamberlain's Method.*—Digest 5 grams of the sample with 250 cc. of 70% (by vol.) alcohol for 24 hours, shaking every half hour during the first 8 hours. Filter through a dry paper, determine nitrogen in 100 cc. of the filtrate by the Kjeldahl or Gunning method, and multiply the result by 5.7. To facilitate evaporation, only 3 cc. of sulphuric acid are first used, the remainder being added after the solution has reached a small volume.

Determination of Salt-soluble Protein.—Chamberlain's Method.*—Digest 12 grams of the flour with 300 cc. of 5% potassium sulphate solution, as described under Alcohol-soluble Protein. Determine nitrogen in 100 cc. of the filtrate, and calculate the salt-soluble protein, using the factor 5.7.

Determination of Acidity of Flour.—Titrate 100 cc. of the solution prepared as described for the determination of nitrites (page 321), with tenth-normal sodium hydroxide solution, using phenolphthalein as indicator. If the distilled water used contains an appreciable amount of carbon dioxide, it should be boiled previous to agitation with the flour.

Determination of Cold-water Extract of Flour.—Wanklyn specified the following method of determining the cold-water extract.

One hundred grams of the flour are thoroughly mixed with distilled water in a graduated liter-flask, which is finally filled with water to the mark; the contents are thoroughly mingled by frequent shaking during six or eight hours, and allowed to stand over night. The supernatant liquid is then poured upon a filter. After rejecting the first few cubic centimeters of the filtrate, exactly 50 cc. are collected and evaporated to dryness in a tared platinum dish on the water-bath. The weight of the dried residue, multiplied by 20, gives the quantity of cold-water extract, which, in a sound flour, according to Wanklyn, should not exceed 5%.

Determination of Iodine Number of the Fat. — Place 20 grams of the flour (or a sufficient quantity to yield 0.2-0.25 grams of fat) in a dish of 100 mm. diameter, and dry in a desiccator over strong sulphuric acid for 3 days, thus removing the larger part of the moisture. Transfer the flour to the inner tube of a Johnson fat extractor, cover with a small amount of cotton previously extracted with ether, weigh down with a

^{*} U. S. Dept. of Agric., Bur. of Chem., Bul. 81, p. 118.

piece of wide-bore glass tubing to prevent the rising of the flour in the extractor, due to expansion of ether in the lower end of the inner tube, and extract for 16 hours with 25 cc. of absolute ether into a tared 35 cc. flask. Drive off the larger part of the ether from the flask by gently heating, and dry to constant weight in the water or steam oven at 100° C., passing a current of dry hydrogen into the flask during the drying to avoid oxidation. As a rule 20 to 30 minutes' drying is sufficient. Reserve the flask and the contents for the determination of the iodine number.

To the 35 cc. flask containing the fat, add 10 cc. of chloroform and 25 cc. of standard Hanus solution. Carefully place the flask, without removal of contents, in a saltmouth glass-stoppered bottle, and let stand one-half hour. Finally break the flask with a glass rod, add 10 cc. of potassium iodide solution and 100 cc. of water, and titrate with standard thiosulphate solution.

Detection of Bleaching in Flour.—The following simple tests serve to determine whether or not a given sample of flour has been bleached with nitrogen peroxide.

Gasoline Test.*—Place 25 grams of the flour in a 4 ounce, wide-mouthed, glass-stoppered bottle, add sufficient gasoline to nearly fill the bottle, shake, and allow to settle. If the flour is unbleached, the gasoline will be distinctly yellow; if bleached, it will remain nearly colorless.

Nitrite Test.—Place 10 grams of the flour in a 4 ounce, wide-mouthed, glass-stoppered bottle. Add 100 cc. of distilled water, and 4 cc. each of sulphanilic acid and alpha-naphtylamine chloride solutions, prepared as directed in the following section. Stopper the bottle and shake vigorously for a few minutes, then allow to stand for at least 30 minutes. If the flour is unbleached, the liquid will not be colored a red tint; if it is bleached, the liquid will take on a color ranging from light pink to deep red, according to the degree of bleaching. Always run for comparison a parallel test on a sample known to be unbleached, so that allowance can be made for any nitrites in the water.

Determination of Nitrites in Flour.—Griess-Ilosvay Method.†—This method, originally designed for the determination of nitrites in water, is well suited for the estimation of the extent to which flour has been bleached by nitrogen peroxide.

^{*} Based on observations of Alway.

[†] Sutton, Volumetric Analysis, 9 Ed., p. 449.

- 1. Reagents.—(a) Sulphanilic acid solution.—Dissolve 0.5 gram of sulphanilic acid in 150 cc. of 20% acetic acid.
- (b) Alpha-naphtylamine chloride solution.—Dissolve 0.2 gram of the salt in 20 cc. of strong acetic acid with the aid of heat. Decant off the clear solution, and make up to 150 cc. with 20% acetic acid.
- (c) Standard sodium nitrite solution.—Prepare pure silver nitrite by mixing a warm concentrated solution of 8 parts of sodium nitrite with a warm concentrated solution of 16 parts of silver nitrate. When cool, wash the precipitate with cold water, and dry quickly in a water bath with as little exposure to the light as possible. Dissolve 0.1097 gram of the dry silver nitrite in warm water, add a slight excess of pure sodium chloride, and make up to 1000 cc. After the silver chloride has settled, draw off 10 cc. of the clear solution, and dilute to one liter. One cc. of this solution contains 0.0001 mg. of the nitrogen as nitrite.
- 2. Determination.—Weigh out 25 grams of the flour (or of the crumbled bread) into an Erlenmeyer flask, add 250 cc. of water free from nitrites, shake vigorously for five minutes, let stand for one-half to one hour, and filter on a paper free from nitrites. Make 50 cc. of the filtrate up to 100 cc. with water, and add 2 cc. each of sulphanilic acid solution and naphtylamine chloride solution, shake, and heat at about 80° C. for 10 minutes to bring out the color.

For comparison, dilute 50 cc. of the standard sodium nitrite solution to 100 cc., and treat with sulphanilic acid and alpha-naphtylamine chloride solutions, as above described. Compare the solutions to be tested with this solution in a colorimeter, and calculate the parts of nitrogen as nitrite per kilo of flour or bread. In flours containing above 3 mgs. per kilo, make 10 cc. (instead of 50 cc.) up to 100 cc., thus avoiding the intense red color obtained in a more concentrated solution.

Bamihl Test for Gluten (Modified by Winton*).—This test serves to detect wheat flour mixed with rye and other flours.

Place a very small quantity of the flour (about 1.5 milligrams) on a microscope slide, add a drop of water containing 0.2 gram of water-soluble eosin in 1000 cc., and mix by means of a cover glass, holding the latter at first in such a manner that it is raised slightly above the slide, and taking care that none of the flour escapes from beneath it. Finally allow the cover glass to rest on the slide, and rub it back and forth until the gluten has collected into rolls. The operation should be carried out on a white paper so that the formation of gluten can be noted.

^{*} U. S. Dept. of Agric., Bur. of Chem., Bul. 122, p. 217.

Wheat flour or other flours containing it yields by this treatment a copious amount of gluten, which absorbs the eosin with avidity, taking on a carmine color. Rye and corn flour yield only a trace of gluten, and buckwheat flour no appreciable amount. The preparations are best examined with the naked eye, thus gaining an idea of the amount of gluten present. Under the microscope traces of gluten, such as are formed in rye flour, are so magnified as to be misleading.

In case the flour is coarse, or contains a considerable amount of branelements, as is true of buckwheat flour and low grade wheat flour, the test should be made after bolting, as the bran particles and coarse lumps interfere with the formation of gluten rolls.

BREAD.

Bread is a term broadly applied to any baked mixture of finely divided grain and water, whether or not other ingredients are used. Pilot, or ship bread, crackers, and unleavened bread, consist almost entirely of flour and water with a slight addition of salt.

Similarly, corn bread or corn cake is frequently made exclusively from corn meal and water. In a narrower sense, however, bread is generally understood to mean the raised or leavened product, rendered light and porous by the aid of gas, which is generated either before or during baking. Commonly the gas employed is carbon dioxide, generated either by the fermentative action of yeast on the sugar of the dough, yielding both alcohol and gas, or by the agency of baking chemicals mixed with the dough, whereby an alkaline bicarbonate is decomposed by the action of an acid to produce the gas. Again, the gas may consist wholly or in part of ammonia, yielded by the vaporization during baking of ammonium carbonate mixed with the dough; and finally, the expansion during baking of the air itself confined in the dough may be the leavening agent, as in the case of puff paste and pie crust.

Wheat flour is of chief value for bread on account of its high content of gluten, in which other cereals are lacking. In the preparation of ordinary white bread, the flour is mixed with water or milk, salt, and yeast, the materials are mingled thoroughly by kneading and allowed to remain for some time in a warm place, during which, by the vinous fermentation induced by the yeast, the mass "rises" or forms a light sponge, due to the action of the gas on the glutinous dough.

During the subsequent process of baking, which should take place at

a temperature between 230° and 260° C., further expansion ensues, much of the water is driven off, and the porous mass sets to form the loaf, the outside of which is converted into a brown crust, due to the caramelizing of the dextrin and sugar into which the starch of the outer layers is converted. Among other changes that take place in the interior or "crumb" during baking are (1) the partial breaking up of the starch grains, which, however, largely retain their identity, though in some degree distorted in shape; (2) somewhat obscure changes in the character of the proteins; and (3) partial oxidation of the oil or fat.

The standard for judging the quality of commercial bread may we'll be based on that of the best home-baked family loaf. The well-made loaf should possess an agreeable odor, and a sweet, nutty flavor, entirely free from mustiness. It should be well "raised," with a good crumbling fracture; it should not be tough or soggy on the one hand (due to underraising), nor extremely dry and spongy on the other (indicative of overraising). Over-raising, moreover, produces sourness, due to advanced lactic fermentation.

Composition of Bread.—The following analyses made in the U. S. Bureau of Chemistry of common varieties of bread were summarized from Bulletin 13, part 9, averages of a number of analyses being given in each case:

	No. of Analyses.	Moisture.	Protein, N×6.25.	Protein, N×5.70.	Ether Extract.
Vienna bread	10	38.71	8.87	8.09	1.06
Home-made bread	2	33.02	7-94	7.24	1.95
Graham bread.	9	34.80	8.93	8.15	2.03
Rye bread	7	33-42	8.63	7.88	0.66
Miscellaneous bread	9 48	34-41	7.60	6.93	1.48
Biscuits or crackers		7.13	10.34	9-43	8.67
Rolls	11	27.98	8.20	7.48	3.41
	Crude Fiber.	Salt.	Ash.	Carbohy- drates, Excluding Fiber.	Calculated Calories of Combus- tion.
Vienna bread.	0.62	0.57	1.19	53-72	4435
Home-made bread	0.24	0.56	1.05	56.75	4467
Graham bread	1.13	0.69	1.59	53.40	4473
Rye bread	0.62	1.00	1.84	56.21	4338
Miscellaneous bread	0.30	0.49	1.00	56.18	4429
	0.47	0.99	1.57	73.17	4755
Biscuits or crackers	0.60	0.60	1.31	59.82	4538

In the examination of bread for its general quality, without regard to its food value, much information may be gained by carefully observing the

physical characteristics of the loaf, its color, taste, odor, porosity, etc. In addition to such data, determination of moisture, ash, and acidity will usually suffice to enable the analyst to pass judgment on its wholesomeness. The following summary gives such analytical data on upwards of fifty samples of bread, purchased from cheaper bakeries and stores, and examined in the author's laboratory.

BREAD.

Kind of Bread.	No. of Analyses.	Weight of Loaf in Grams.	Water, Per Cent.	Per Cent Ash in Terms of Solids.	Acidity.*
White	44				
Maximum		653	45.20	1.83	6.2
Minimum		126	33.00	0.60	1.3
Mean	••	430	40.72	0.85	2.6
Graham	7			·	
Maximum	••	500	45.20	1.55	4.2
Minimum	••	367	40.10	0.96	2.1
Mean	••	420	41.50	1.26	3.5
Whole wheat	I	507	45.10	1.20	
Diabetic	I	445	47.00	2.20	
Muffins	I	194	48.20	1.15	1.7
Rye	I	1291	47.15	2.13	10.0
'Black''	I	550	47.00	2.20	
German with seeds	I	417	42.30	0.95	
Brown	I	500	48.10	3.50	
'Knackerbrod"	I	110	8.00	1.94	

^{*} Cubic centimeters of tenth-normal soda required to neutralize 10 grams of the fresh bread.

Water in Bread.—The amount of water is of considerable importance, and, in the best bread, varies from 33 to 40 per cent. A larger content of water than 40% should be considered objectionable in a white bread, both on the ground of acting as a make weight, and because a large excess of moisture tends to cause the growth of mold.

Acidity of Bread.—The degree of sourness of a sample of bread is one of the most important indications as to its quality, and is most readily obtained by rubbing up in water, by means of a pestle, 10 grams of the "crumb," and titrating with tenth-normal alkali, using phenolphthalein as an indicator. To neutralize the acidity of 10 grams of the normally sweet loaf, an average of 2 cc. of the standard alkali solution is required, corresponding to 0.72 gram of lactic acid per loaf of an average weight of 400 grams. The loaf exhibiting the maximum sourness or acidity in the above table required 10 cc. of standard alkali per 10 grams of bread, corresponding to 11.61 grams lactic acid in the loaf of 1,201 grams.

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Fat in Bread.—It is well known that the results of fat or ether extract as obtained by the ordinary method and expressed in most bread analyses are too low, being considerably less than the combined fat of the materials entering into its composition. This is probably due to the fact that during baking the fat particles are incrusted with insoluble matter, which protects them from the subsequent action of the ether. It is further claimed by some that the partial oxidation of the fat during baking has something to do with the low results. No perfectly satisfactory improvement over the regular ether method for fat extraction in bread has been discovered, and therefore this method, as described elsewhere, is recommended.

Adulteration of Bread.—The fraudulent addition of inert foreign ingredients to bread is almost never practiced, and is mainly of historic interest. Gypsum, chalk, bone ash, and various other minerals have been mentioned as possible adulterants, but the amount of any of these materials necessary to add for purposes of profit could scarcely be present without very apparent injury to the quality of the bread. Their presence in any considerable degree would be apparent in the abnormally high ash content of the bread.

The employment of alum to "improve" inferior or unsound flour has already been referred to, and, for the same purpose, sulphate of copper in small quantities is also said to have been used, enabling the making of bread of fairly good appearance from flour that was distinctly damaged.

Alum in Bread * is tested for by a modification of the logwood process described on page 315 as follows: 5 cc. of the logwood tincture and 5 cc. of the saturated ammonium carbonate solution are diluted to 100 cc., and the freshly prepared mixture poured over about 10 grams of the bread crumbs in a porcelain evaporating-dish. After standing a few minutes, as much as possible of the liquid is drained off, the bread is slightly washed by one treatment with water, and dried in the water-oven. In presence of alum, a dark-blue color is given to the bread, which becomes deeper on drying. The color is proportional to the amount of alum present. If the sample is free from alum, the color varies from red to light brown. The reagent solution must be freshly prepared. This test is not perfectly reliable in the case of very old or sour breads, which have been known to give the color test with logwood in the absence of alum.

^{*} Jago on Bread, p. 634.

Copper Salts in Bread are detected in the ash by the same method as that used for canned goods (p. 902).

Cake and Similar Preparations.—These differ from bread chiefly by the addition of considerable sugar, butter, spices, and other flavoring materials. In gingerbread, molasses is used as an important ingredient besides ginger. The adulterants of molasses, such as glucose, salts of tin, etc., would thus sometimes occur in gingerbread. In fact stannous chloride has been found in ginger cakes.*

The following analyses of a few typical varieties of cakes are selected from Bulletin 13 of the Bureau of Chemistry:

	Moisture.	Proteins, N×6.25.	Proteins, N×5.70.	Ether Extract.	Crude Fiber.
Doughnuts	21.61	6.73	6.14	19.33	0.60
Ginger snaps	4.86	6.06	5-53	15.44	0.79
Fruit cake	24-47	4.56	4.16	12.35	1 .,
Gingerbread	21.49	6.25	5.70	8.42	0.90
Cup cakes	14.81	5.24	4.78	15.56	0.27
Macaroons	8.06	6.67	6.08	12.97	1.41
Jumbles	13.34	7.62	6.95	14.79	1.04
				 C1-1	<u> </u>
	Ash.	Salt.	Sugar.	Carbohy- drates other than Fiber and Sugar.	Calculated Calories.
Doughnuts			Sugar.	drates other than Fiber and Sugar.	Calories.
Doughnuts	O.40	0.03 0.47		drates other than Fiber and	
Ginger snaps	0.40	0.03	1.28	drates other than Fiber and Sugar.	Calories.
Ginger snaps	0.40	0.03	1.28 28.66	drates other than Fiber and Sugar. 50.64 24.90	5529 4971 4757
Ginger snaps	0.40 1.82	0.03 0.47 0.28	1.28 28.66 9.48	drates other than Fiber and Sugar. 50.64 24.90	5529 4971

YEAST.

The yeast plant is a fungus of the genus Saccharomyces, widely distributed through the vegetable kingdom and in the air. It is capable of rapid growth by the multiplication of its cells when present in a favorable medium, such as malt wort, and with propitious conditions of temperature, moisture, etc. Under such conditions, it forms a yellowish, viscous, frothy substance, the chief value of which, in the liquor industry, is the production of alcohol, while for bread-making, as a result of the same kind of fermentation, the end desired is the leavening of the doughy mass by the carbon dioxide liberated.

^{*} See U. S. Dept. of Agric., Bur. of Chem., Bul. 13, p. 1369.

A vigorous, pure yeast which will "raise" quickly is a great preventive against sour bread, for not only is it comparatively free from the germs and products of lactic acid fermentation, but by doing its work quickly it enables the baker to check the fermentation or raising process before the lactic acid or sour decomposition is far advanced.

Yeast most commonly used in bread-making is of the so-called "compressed" variety. The use of compressed yeast is almost universal for domestic purposes, and is more or less common in bakeries. A small amount of brewers' yeast in liquid form from beer wort is used, especially in the immediate neighborhood of breweries, and dry yeasts are used to some extent in localities so remote that fresh compressed yeast cannot readily be obtained.

Compressed Yeast is a product of distilleries where malt and raw grain are fermented for spirits. Most of it comes from whisky wort, and some from the worts used in the manufacture of gin and other distilled liquors. Little if any of the commercial compressed yeast is made from beer wort yeast.

In the manufacture of compressed yeast, the yeast floating on the top of the wort is separated by skimming, while that settling to the bottom is removed by running the wort into shallow settling trays. Top yeast is considered more desirable than bottom yeast for bread-making. The separated yeast is washed in cold water, and impurities are removed, either by sieving through silk or wire sieves, or by fractional precipitation while washing. The yeast, with or without the addition of starch, is finally pressed in bags in hydraulic presses, after which it is cut into cakes, packed in tin-foil, and kept in cold storage till distributed for use.

Such yeast should be used when fresh, as it readily decomposes and soon becomes stale. When fresh, it should have a creamy, white color, uniform throughout, and should possess a fine, even texture; it should be moist without being slimy. It should quickly melt in the mouth without an acid taste. Its odor is characteristic, and should be somewhat suggestive of the apple. It should never be "cheesy," such an odor indicating incipient decomposition, as does a dark or streaked color.

Dry Yeast is prepared by mixing fresh yeast with starch or meal, molding into a stiff dough, and drying, either in the sun or at a moderate temperature under reduced pressure. Such yeast, when dry, is cut into cakes and put in packages. It will keep almost indefinitely. During the drying process, many of the yeast cells are rendered torpid and temporarily inert, and for this reason the dried yeast does not act so promptly

in leavening as does compressed or brewers' yeast, but when once it begins to act it is quite as efficacious.

Composition of Yeast.—The following is the result of the analysis of under-fermentation yeast, after drying, by Nägele and Loew:

Cellulose and mucilage	37
Albuminoids (mycroprotein, etc.)	36
" soluble in alcohol	9
Peptones (precipitable by subacetate of lead)	2
Fat	5
Extractive matters (leucin, glycerin, etc.)	4
Ash	7
•	100

Lintner gives the following average analyses of the ash of three samples of yeast, analyzed by him:

Silica	1.34
Iron (Fe ₂ O ₃)	0.50
Lime (CaO)	5-47
Sulphuric anhydride (SO ₈)	0.56
Magnesin (MgO)	•
Phosphoric anhydride (P ₂ O ₅)	
Potash (K ₂ O) and a little soda	•
	08.08

Matthews and Scott give the following as the ash composition of yeast:

Potassium phosphate	78.5
Magnesium phosphate	13.3
Calcium phosphate	6.8
Silica, alumina, etc	1.4
•	
	100.0

Microscopical Examination of Yeast.—Mix a bit of the yeast in water on the glass slip till a milky fluid is formed, and stir in a drop of a very weak anilin dye solution, such as methyl violet, eosin, or fuchsin.* Put on the cover-glass, and examine under the microscope. Living, active cells resist the stain, if the latter is dilute enough, and appear colorless or nearly so, while the decayed and lifeless cells are stained, and can easily be distinguished by their color. Yeast cells are circular or oval in shape, and vary from 0.007 to 0.009 mm. in diameter. They are sometimes isolated, and sometimes grouped in colonies; each cell has an outer, mucilaginous coating or envelope. The interior, granular mass or substance of the cell is the protoplasm, and within the protoplasm are frequently seen one or more circular empty spaces known as vacuoles.

Yeast cells multiply by the process of budding. The decadence of yeast cells is marked by the increased size of vacuole, and by the thickening of the cell wall.



Fig. 68.—Sprouting Yeast-cells (Saccharomyces cerevisia). (a, after Lürssen; b, after Hansen.)

Yeast-testing.—Available Carbon Dioxide.—The value of yeast in bread-making depends on the amount of carbon dioxide which it is capable of generating under given circumstances, hence the available carbon dioxide is the chief factor in gauging a yeast. There are various methods of determination, (1) either by measuring the volume of gas set free by the action of a weighed quantity of yeast in a sugar solution of known strength, kept for a fixed time at a fixed temperature (say 30°), or (2) by conducting the gas from such a fermenting solution through a weighed absorption bulb, containing potassium hydroxide and noting the increase in weight, or (3) by the more convenient method of Meissl as follows:

A mixture is made of 400 grams pure, concentrated sugar, 25 grams ammonium phosphate, and 25 grams potassium phosphate. A small, wide-mouthed flask of about 100 cc. capacity is fitted with a doubly perforated rubber stopper, having two tubes as shown, one of which is bent and passes nearly to the bottom of the flask, being fitted at the outer end with a rubber tube and glass plug, while the other is connected with a small calcium chloride tube. Measure 50 cc. of distilled water into

^{* 1} gram crystallized fuchsin in 160 cc. water having 1 cc. alcohol.

this flask, and dissolve 4.5 grams of the above sugar phosphate mixture. Finally add 1 gram of the yeast to be tested, stir it well till there are no

lumps, and cork the flask. Carefully weigh on a delicate balance the flask with its contents, and immerse in a water-bath at 30° C., keeping it at that temperature for six hours. At the end of this time, remove the flask from the bath, and immediately immerse in cold water to cool the contents. Remove the rubber tube with the glass plug, and by suction draw out the remaining carbon dioxide. Replace the plug, and having carefully wiped off the flask, again weigh. The loss in weight is due to carbon dioxide set free by the fermentation of the yeast.

Starch in Compressed Yeast.—The addition of potato starch to yeast before pressing has long been customary, on the grounds that the starch acted as a drier, producing a much cleaner product, and one that could be more readily and intimately mingled with the materials of the bread, besides enhancing the keeping qualities of the yeast, es-



Fig. 69.—Apparatus for Determining Leavening Power of Yeast.

pecially in warm weather. The best grades of compressed yeast contain about 5% of starch, but some are found with 50% and even more. Undoubtedly the larger amounts are added as a make weight.

The question has frequently been raised whether, with improved methods of manufacture, whereby yeast could be produced comparatively free from slime, and thus capable of pressure without the admixture of starch, the use of the latter should not be considered as an adulterant. Some of the compressed yeast on the market is free from starch, and its makers claim that this is the only absolutely pure variety, while the presence of starch should be distinctly regarded as a violation of the section of the food law which forbids the use of a cheaper or inferior ingredient.

Briant claims that the admixture of starch up to 5% increases rather than decreases the actual content of yeast, in that the starch abstracts moisture from the yeast cells themselves, the proportion of water being much smaller, and that of the yeast larger in the starch-mixed substance. T. J. Bryan,* on the other hand, finds that the addition of starch to yeast reduces the carbon dioxide value, and that the percentage reduction is greater than the percentage of starch present. His experiments further

^{*} A. O. A. C. Proc. 1907, U. S. Dept. of Agric., Bur. of Chem., Bul. 116, p. 25.

indicate that the keeping qualities of starch yeast is not greater but actually less than that of pure yeast.

In the absence of a legal standard for starch in yeast, it is difficult to see how complaints could be maintained under the general food laws of most states, without condemning the use of starch altogether.

Jago suggests 20% as the limit for starch in yeast, beyond which it should be considered as an adulterant.

CHEMICAL LEAVENING MATERIALS.

Under this heading are included the various ingredients that enter into the mixtures commonly known as "baking powders" which have no food value in themselves, but are, strictly speaking, instruments or tools that by purely chemical reactions bring about, under certain conditions, the comparatively quick liberation of gas and the consequent aeration of biscuit, bread, and cake.

Baking Powders and their Classification.—Formerly the housewife was accustomed to measure out in proper proportion a mixture of sour milk, or cream of tartar, with saleratus to produce quick aeration of bread. The modern baking powder is a natural outgrowth of the former practice, and has almost wholly displaced it, producing, as it does, a mixture ready for immediate use of an acid and an alkaline constituent in proper proportion for chemical combination to form the gas. A third ingredient is, however, generally considered as necessary to check deterioration, viz., a dry, inert material, which by absorbing moisture prevents the premature chemical action between the reagents. Starch is nearly always used for this purpose, though sugar of milk has a limited use. The alkaline principle of nearly all baking powders is bicarbonate of soda, or saleratus. Baking powders are divided naturally into three main classes, with reference to the acid principle:

(1) Tartrate Powders, wherein the acid principle is (a) bitartrate of potassium or (b) tartaric acid, typified by the following reactions:

(2) Phosphate Powders, in which calcium acid phosphate is the acid principle:

(3) "Alum Powders," wherein the acidity is due wholly or in part to sulphate of aluminum as it occurs in potash or ammonia alum, or in the double sulphates of aluminum and sodium.*

Assuming burnt potash alum as the substance used, the reaction would be as follows:

Naturally many baking powders of complex composition are met with, embodying various mixtures of the above classes.

Composition of Various Baking Powders.—Following are analyses of typical baking powders of the above classes: †

1. Cream of Tartar Baking Powder:

Total carbon dioxide, CO ₂	13.21
Sodium oxide, Na ₂ O	13.58
Potassium oxide, K ₂ O	14.93
Calcium oxide, CaO	.18
Tartaric acid, C ₄ H ₄ O ₅	41.60
Sulphuric acid, SO ₃	.10
Starch	7-42
Water of combination and association by difference	8.98

Available carbon dioxide 12.58%.

100.00

^{*}It is probable that very little ammonia or potash alum is actually used at present in this class of powders. A product largely used is known in the trade as C. T. S. (cream of tartar substitute) and is a calcined double sulphate of aluminum and sodium.

[†] Div. of Chem., Bul. 13, part 5, pp. 600, 604, and 606.

2. Phosphate Baking Powder:	
Total carbon dioxide, CO ₂	13.47
Sodium oxide, Na,O	12.66
Potassium oxide, K ₂ O	.31
Calcium oxide, CaO	10.27
Phosphoric acid, P,O,	21.83
Starch.	26.41
Water of combination and association by difference	15.05
-	100.00
Available carbon dioxide 12.86%.	
Alon Palina Dandon	
3. Alum Baking Powder:	
Total carbon dioxide, CO ₂	9-45
Sodium oxide, Na ₂ O	9.52
Aluminum oxide, Al ₂ O ₃	3.73
Ammonia, NH ₃	1.07
Sulphuric acid, SO ₃	10.71
Starch.	43.25
Water of combination and association by difference	22.27
	100.00
Available carbon dioxide 8.10%.	
Mixed Powders:	
Total carbon dioxide, CO ₂	10.68
Sodium oxide, Na ₂ O	14.04
Calcium oxide, CaO.	1.29
Aluminum oxide, Al ₂ O ₃	4.59
Ammonia, NH ₃	1.13
Phosphoric acid, P ₂ O ₅	3.38
Sulphuric acid, SO ₃	11.57
Starch.	42.93
Water of combination and association by difference	10.39
	100.00

Available carbon dioxide 10.37%.

The Adulteration of Baking Powder.—No substance that comes within the domain of food inspection is the subject of so much controversy

as baking powder. Unless a specific law forbids the use of a particular ingredient or class of ingredients, or in some manner regulates the labelling of the package, no baking powder of any kind can be considered adulterated under the general food law, unless it can be proved to be injurious to health, or unless it contain inert and useless mineral matter.

As a matter of fact, the residue left in the bread by all classes of baking powder consists of one or more drugs recognized in the Pharmacopæia. all of which in large quantity exercise well-marked toxic effects on the human system. Artificial digestion experiments, and physiological tests on the lower animals, using excessive doses of any of the above drugs, do not show the effect of the every-day use of baking powder in bread on the human system, and only a systematic examination of the effect of such use on large numbers of people can prove conclusively whether or not any one class of baking powders is harmful, and hence whether or not it should be classed as adulterated. Aside from the question of the harmfulness of the acid ingredients employed in baking powder, which is the subject of much controversy among rival manufacturers, there can be no doubt that such inert make weight substances as calcium sulphate. or terra alba, or clay, which are entirely useless, and lower the strength, quality, and purity of the powder, are to be considered in the light of adulterants.

Cream of Tartar—Its Nature and Adulteration.—Cream of tartar, or potassium bitartrate (KH₆C₄O₆), is the purified product obtained by the recrystallization of the crude argols or lees deposited in the interior of wine casks.

The lees, or argols, consist chiefly of crude potassium bitartrate, which is present in the juice of the grape, but is insoluble in the alcohol formed in the fermentation, and is hence deposited. If, for the clarification of the wine, such substances as gypsum or plaster of Paris are used, tartrate of calcium will be found mixed with the bitartrate of potassium in the lees. Hence it is that calcium tartrate is sometimes found in commercial cream of tartar.

Potassium bitartrate is insoluble in alcohol, sparingly soluble in cold, and readily soluble in hot water.

Allen * states that when the calcium tartrate is present in excess of 10%, it should undoubtedly be considered as an adulterant.

Other common adulterants of cream of tartar are calcium acid phosphate, gypsum, or plaster of Paris, starch, and alum.

CHEMICAL ANALYSIS OF BAKING CHEMICALS AND BAKING POWDERS.

Cream of Tartar.—The degree of purity of commercial cream of tartar is best determined by weighing out exactly 0.188 gram of the sample, dissolving in hot water, and titrating with tenth-normal sodium hydroxide, using phenolphthalein as an indicator. If the article is pure, exactly 10 cc. of the standard alkali will be required for the titration. All the above-named adulterants, with the exception of alum, are either insoluble, or sparingly soluble in hot water, and will indicate the impurity of the sample even before titration. If the adulterant be alum, the sample would go into solution in the water, but the alum would be precipitated by the sodium hydroxide, the precipitate being, however, soluble in an excess of the alkali.

Sodium Bicarbonate on account of its cheapness is rarely adulterated, save by the occasional presence of common salt, an impurity incidental to its manufacture. The degree of purity of sodium bicarbonate is best ascertained by titration with standard acid, each cubic centimeter of tenthnormal acid being equivalent to 0.0084 gram of sodium bicarbonate.

Determination of Total Carbon Dioxide.—Reagents.—Calcium Chloride.—This can be obtained in granulated form in pellets of about the size of peas, specially prepared for moisture absorption.

Soda Lime.*—To a kilogram of commercial sodium hydroxide, 500 to 600 cc. of water are added, and the mixture heated in an iron kettle to form a thin paste. While still hot, a kilogram of coarsely powdered quick-lime is added, stirring with an iron rod. The lime is slaked, and the whole mass heats and steams up. No outside heat is necessary at this stage, but the mass is stirred and the lumps broken up. As soon as cool, place the product in wide-mouthed bottles, and seal with paraffin wax. The product should be slightly moist to give the best results.

Hydrochloric Acid.—Specific gravity 1.1.

Sulphuric Acid.—Specific gravity 1.85.

Potassium Hydroxide Solution.—Specific gravity 1.55.

Two varieties of apparatus are in use for the determination of carbon dioxide. In one form the amount of carbon dioxide is obtained by difference in weight of the apparatus, before and after elimination of the gas. In the other, the gas driven out of a given weight of the sample is absorbed, and its amount calculated from the increase in weight of the

^{*} Benedict and Tower, Jour. Am. Chem. Soc., Vol. XXI, p. 396.

Types of these varieties are the Geissler and the Knorr absorbent. apparatus

The Geissler Apparatus.—This consists of a flask A, having a ground neck a, and a flaring funnel-top A'. B is an elongated bulb, closed at the top by the hollow stopper K, and terminating below in the hollow stem B', which is accurately ground at b to fit the neck a. Fused into the bulb B is the tube C, and within this is the small tube D, open at the top and communicating directly with the hollow stem B'. gg are openings between B and C.

E is a fine glass tube, passing from the bottom of the hollow stem B' and to the height of a small protuberance e in the bottom of the funnel A', the construction being such that by turning the bulb and stem BB' in the neck a of the flask A the tube E may be opened or closed at the top. His a side tube in the flask A, closed by the ground stopper h.

The bulb B and the tube C are filled with strong sulphuric acid nearly to the top of the tube D, by passing through the neck at the top, which is then closed by the stopper K.

About 0.5 gram of the dried sodium bicarbonate, or I gram of the baking powder, is introduced into the flask A through the neck a from a weighing-tube or otherwise, so that its exact weight is known. The stem B' is then inserted, and the funnel-top A' is nearly filled with the hydrochloric acid, the tube e being Fig. 70,-Geissler's CO, Apclosed.

paratus or Alkalimeter.

The entire apparatus is then weighed, after which the stem is turned to bring the protuberance e nearly opposite the tube E, uncovering it enough to allow the acid to pass slowly down the tube into the flask and upon the powder in the bottom of the flask. The carbon dioxide evolved passes through the opening f into the hollow stem B', thence up through the tube D, and down and up (as indicated by the arrows) through the sulphuric acid, which absorbs the moisture. Finally the gas passes out through the tube K.

After the evolution of the gas has continued for two or three minutes, gentle heat is applied to the flask from a gas flame, and the solution is brought to boiling, which is continued for a few minutes, during the latter portion of which the stopper h is removed, and the tubulure connected by rubber tubing with a system of two U tubes, one containing soda lime, and the other calcium chloride. The tube k is then connected with the aspirator, and a current of dried air is passed through the apparatus at the rate of about two bubbles per second, long enough to displace all the carbon dioxide. The rubber tubes are then disconnected, the stopper K is replaced, and the apparatus cooled to room temperature and weighed.

The available carbon dioxide in baking powder is determined in the same manner as above, by simply substituting freshly boiled, distilled water for the hydrochloric acid in the funnel-top A'.

The Knorr Apparatus (Modified).—The apparatus (Fig. 71) consists of (1) a flask, into which is introduced an accurately weighed amount of

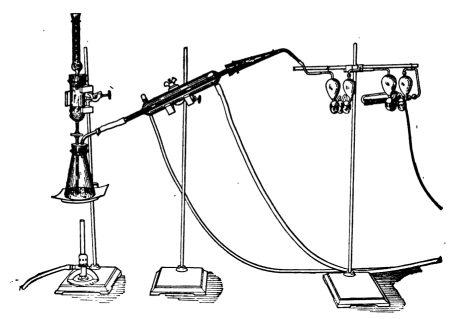


Fig. 71.-Modified Knorr Apparatus for Determining Carbon Dioxide.

the dry sample (0.5 to 1 gram of sedium bicarbonate or 1 to 2 grams of baking powder); (2) a funnel, the tube of which, provided with a stop-cock enters the stopper of the flask; (3) a soda lime tube, entering a stopper at the top of the funnel; (4) a Liebig condenser, connecting with a tube passing through the stopper of the flask; (5) a Geissler bulb, filled with the sulphuric acid; (6) a potash absorption-bulb, and (7) a calcium

chloride tube, which may if desired be replaced by a second sulphuric acid bulb. The potash absorption apparatus is accurately weighed before being connected up, and the funnel is nearly filled with the hydrochloric acid reagent, after which the soda lime tube is attached. The calcium chloride tube is connected by a rubber tube with the aspirator, and a current of cold water is allowed to run through the outer Liebig condenser-tube.

The stop-cock in the funnel-tube is first opened to allow the acid to slowly run into the flask, the flow being regulated to insure slow evolution of the gas.

The aspirator is then turned on so that about two bubbles of air per second pass through the apparatus, and gentle heat is applied to the flask by the gas flame, the solution within being brought to boiling, and the boiling continued for several minutes after the vapor has begun to gather in the condenser.

Prolonged boiling of the solution should be avoided, and in a series of tests the time of boiling should be precisely the same in all cases.

After removing the flame, the flask is allowed to cool, the aspiration being continued. The absorption-tube is then removed and weighed at room temperature, the increase in weight being due to the carbon dioxide.

The Available Carbonic Acid in Baking Powder is determined in the same manner as the total carbon dioxide, except that recently boiled, distilled water is substituted for the hydrochloric acid.

Detection of Tartaric Acid.*—It is often desirable to test a "compound" cream of tartar, or a "cream of tartar substitute," or an adulterated sample made up largely of foreign ingredients, to see if any tartaric acid, free or combined, be present. The following test is applicable in presence of phosphates:

If the substance to be tested is found to be free from starch, mix a little of the dry powder in a test-tube with a bit of dry resorcin, add a few drops of concentrated sulphuric acid, and heat slowly. A rose-red color indicates tartaric acid or a tartrate, the color being discharged on dilution with water.

In case of baking powder, or a cream of tartar substitute containing starch, shake repeatedly from 3 to 5 grams of the sample with about

^{*} Wolff, Rev. Chim. Analyt. et appr. 4 (1899), p. 2631.

250 cc. of cold water in a large flask, allowing the insoluble portion to subside.

Decant the solution through a filter, and evaporate the filtrate to dryness, after which test the dried residue or a portion thereof with resorcin and sulphuric acid as above described.

Determination of Total Tartaric Acid. — Modified Heidenhain Method.*—Applicable only in the absence of phosphates and salts of aluminum and calcium.

Into a shallow porcelain dish, 6 inches in diameter, weigh out 2 grams of the material and sufficient potassium carbonate to combine with all tartaric acid not in the form of potassium bitartrate. Mix thoroughly with 15 cc. of cold water, and add 5 cc. of 99% acetic acid. Stir for half a minute with a glass rod bent near the end. Add 100 cc. of 95% alcohol, stir violently for five minutes, and allow to settle at least thirty minutes. Filter on a Gooch crucible with a thin layer of paper pulp, and wash with 95% alcohol until 2 cc. of the filtrate do not change the color of litmus tincture diluted with water. Place the precipitate in a small casserole, dissolve in 50 cc. of hot water, and add standard fifth-normal potassium hydroxide solution, leaving it still strongly acid. Boil for one minute. Finish the titration, using phenolphthalein as indicator, and correct the reading by adding 0.2 cc. One cc. of fifth-normal potassium hydroxide solution is equivalent to 0.026406 gram tartaric anhydride (C,H,O,), 0.03001 gram tartaric acid (H,C,H,O_a), and 0.03763 gram potassium bitartrate (KHC,H,O₆).

The standard of the potassium hydroxide solution should be fixed by pure dry potassium bitartrate.

The accuracy of this method is indicated by the agreement of the percentages of potassium bitartrate in cream of tartar powders containing no free tartaric acid, obtained by calculation from the tartaric acid, with those obtained by calculation from the potassium oxide.

In presence of phosphates or of aluminum and calcium salts, the only satisfactory method of arriving at the amount of tartaric acid present is by difference, having determined or calculated the other ingredients.

Kenrick's Polariscopic Methods.—Method 1. (Applicable to Cream of Tartar).—The method is based on the fact that in the presence of excess of ammonia, the rotation of the solution is proportional to the

^{*} Provisional methods of the A. O. A. C., Bur. of Chem., Bul. 65, p. 104; Bul. 107 (rev.), p. 175.

concentration of the tartaric acid, and is independent of the other bases and acids present.

(a) The Substance is Completely Soluble in Dilute Ammonia.—A weighed quantity of the material containing not more than 1 gram tartaric acid is placed in a 25 cc. measuring flask, moistened with 3 or 4 cc. of water, and concentrated ammonia (sp. gr. 0.880) added in quantity sufficient to neutralize all acids that may be present, and leave about 1 cc. in excess. The actual amount of the excess is not of importance, but a greater quantity than 1 cc. of free ammonia should be avoided. The solution is then made up to 25 cc. with water, filtered, if necessary, through a dry filter, and measured in a 20 cm. tube in the polarimeter.

The amount of tartaric acid (C₄H₆O₆) in grams (y) in the material taken is given by the formula:

$$y = 0.00519x$$

where x is the rotation in minutes.

(b) The Substance is not Completely Soluble in Dilute Ammonia.—In this case calcium tartrate is probably present, and may be determined as follows: Treat I gram of the substance (or an amount containing not more than I gram of tartaric acid) in a small beaker with 15 cc. of water, and 10 drops of concentrated hydrochloric acid. Heat gently till both the potassium and calcium tartrates have passed into solution, and then, while still hot, add 2 cc. of concentrated ammonia (or enough to produce an ammoniacal smelling liquid), and about 0.1 gram of sodium phosphate dissolved in a little water. Transfer to a 25-cc. measuring flask, cool, make up to the mark with water, filter through a dry filter, and polarize the filtrate in a 20-cm. tube. The tartaric acid is calculated from the formula given under (a).

The precipitation of the calcium by means of sodium phosphate is not absolutely necessary, but when this is not done, in cases where the proportion of calcium in the sample is high, there is a great tendency for the calcium tartrate to crystallize out from the ammoniacal solution before the reading is made.

The tartaric acid present as bitartrate of potash may be determined by proceeding as in (a), the calcium tartrate being practically insoluble in cold ammonia solution.

The tartaric acid present as calcium tartrate is given, with sufficient accuracy for most purposes, by the difference between the results of (a) and (b). If more accurate results are required, the residue insoluble in

ammonia in (a) may be dissolved in a little hydrochloric acid and treated as above with sodium phosphate and ammonia.

Method 2. (Applicable to Baking Powder and Cream of Tartar mixed with Substitutes).—Direct readings of rotation in ammoniacal solution are inadmissible in analyses of the substances of this class, on account of the influence of iron and aluminum on the rotation of tartaric acid, and on account of the small but unknown rotation of the trace of inverted starch.

Accurate determinations, however, may be made in the presence of excess of ammonium molybdate in neutral solution. The latter substance has the property of greatly increasing the rotation of tartaric acid, so that by its use the small rotation of the inverted starch is made insignificant. It is to be noted, however, that this increased rotation is very sensitive to the presence of alkali and acid, and is, moreover, modified by phosphates. It is therefore necessary, in the first place, to remove the phosphoric acid, and, secondly, to bring the solution to a definite state of neutrality. Both these results are attained by the following procedure, the details of which must be carefully adhered to:

(a) Reagents.—The following solutions must be prepared, but need not be made up very accurately:

Molybdate solution: 44 grams ammonium heptamolybdate in 250 cc. Citric acid solution: 50 grams citric acid in 500 cc.

Magnesium sulphate solution: 60 grams MgSO₄. 7H₂O in 500 cc.

Ammonia solution: 80 cc. concentrated ammonia (sp. gr. 0.880) in 500 cc.

Hydrochloric acid: 60 cc. concentrated hydrochloric acid in 500 cc. Methyl orange solution:

(b) Process.—An amount of material containing not more than 0.2 gram tatraric acid, not more than 0.3 gram alum, and not more than 0.3 gram calcium superphosphate, is accurately weighed, and placed in a dry flask. To this, 5 cc. of citric acid and 10 cc. of molybdate solution are added, and allowed to react with the substance for 10 or 15 minutes (with an occasional shake). Next, 5 cc. of magnesium sulphate solution are added, and 15 cc. of ammonia solution stirred in. After a few minutes (not more than one hour), the solution is filtered through a dry filter, a slight turbidity of the filtrate being disregarded. To 20 cc. of the filtrate are then added a few drops of methyl orange and hydrochloric acid, from a burette, till the pink color appears (2 or 3 drops too much or too little are of no consequence). Finally, 10 cc. more of the molybdate

solution are added to the pink solution, which now becomes colorless or pale yellow, and water is added to make up the volume to 50 cc. This solution, after filtering if necessary, is polarized in a 20-cm. tube.

The amount of tartaric acid in grams (y) in the weight of substance originally taken is given by the following formula, in which x is the rotation in minutes:

$$y = 0.001086x + 0.001601\sqrt{x}$$

But if the rotation is not less than 40', the simpler formula;

$$y = 0.0075 + 0.001168x$$

may be employed.

The following table gives the tartaric acid in grams for every 10 minutes rotation:

Rotation in Minutes.	Grams Tartaric Acid.	Rotation in Minutes.	Grams Tartaric Acid.
10	0.016 0.029 0.0415 0.0535 0.0657 0.0776 0.0895	90	0.1130 0.1246 0.1365 0.1479 0.1595 0.1710 0.1825

Determination of Starch.—McGill's Method* (Modified).—Digest 1 gram of the sample with 150 cc. of a cold 3% solution of hydrochloric acid during twenty-four hours, with occasional shaking. Filter through a tared Gooch crucible, wash first with water until neutral, then once with alcohol, and finally with ether. Dry at 110° C. for four hours, cool, and weigh. Burn off the starch, and again weigh. The difference in the two weights indicates the weight of the starch. The purity of the starch is insured by examination with the microscope.

Acid Conversion Method.†—If the sample contains lime, mix 5 grams in a 500-cc. flask with 200 cc. of 3% hydrochloric acid, and let the mixture stand an hour with frequent shaking. Filter through a wetted 11-cm.

^{*} Canada Inland Rev. Bul. 68, p. 33.

[†] U. S. Dept. of Agric., Bur. of Chem., Bul. 65, p. 105; Bul. 107 rev., p. 176.

filter, wash with water, and transfer the starch by a wash-bottle from the filter-paper back into the original flask, using 200 cc. of water.

If the sample be free from lime, weigh 5 grams directly into the 500-cc. flask with 200 cc. of water. In either case add 20 cc. of hydrochloric acid (specific gravity 1.125) and heat the flask in boiling water for $2\frac{1}{2}$ hours, the flask being provided with a reflux condenser. Determine the dextrose, and from this the starch in the regular manner.

Detection of Aluminum Salts.*—(a) In Baking Powder.—Applicable in presence of phosphates. Burn to an ash about 2 grams of the sample in a platinum dish. Extract with boiling water and filter. Add to the filtrate sufficient ammonium chloride solution to produce a distinct odor of ammonia. A flocculent precipitate indicates aluminum.

In igniting, as above directed, sodium aluminate results from the more or less complete fusion. The reaction which occurs may be represented as follows:

If any phosphate of lime be present, it will be insoluble in the solution of the ash. If phosphate of sodium or potassium be present, it will go into solution, but will only precipitate out when an aluminum salt is also present on the addition of the ammonium chloride reagent.

(b) In Cream of Tartar.—Mix about I gram of the sample with an equal quantity of sodium carbonate, burn to an ash, and proceed as in the case of baking powder (a).

Determination of Alumina.—The above qualitative method with ammonium chloride may be made quantitative in presence of phosphates as follows: After carrying out the qualitative method as above directed, filter off the final precipitate, dissolve it in nitric acid, and test it for phosphate with ammonium molybdate. If phosphates are found absent, proceed as before with a weighed amount of the sample and wash, ignite, and weigh the residue as Al₂O₃.

If phosphate is found present in the ammonium chloride precipitate, proceed as before, igniting and weighing the total residue. Then determine the P_2O_5 in the latter and subtract from the total. The difference will be the Al_2O_3 .

^{*}Leach, 31st An. Rep. Mass. State Board of Health, 1899, p. 638.

Determination of Lime.—5 grams of the sample are treated in a 500-cc. graduated flask with 50 cc. of water and 25 cc. of concentrated hydrochloric acid. Add water to the mark, shake, and allow the starch to settle. Decant through a dry filter, and to 50 cc. of the filtrate add ammonia nearly to neutralization, an excess of ammonium acetate solution, and 4 cc. of 80% acetic acid, and heat at 50° C. Filter if necessary, and precipitate the lime with an excess of ammonium oxalate. Filter, wash, and ignite over a blast-lamp. Weigh as CaO.

Determination of Potash and Soda.*—Weigh out 5 grams into a platinum dish, and incinerate in a muffle at a low heat. The charred mass is well rubbed up in a mortar, then boiled fifteen minutes with about 200 cc. of water, to which has been added a little hydrochloric acid. The whole is transferred to a 500-cc. flask, and, after cooling, made up to the mark and filtered. Of the filtered liquid 100 cc., representing 1 gram of the sample, are measured out, heated to boiling, and a slight excess of barium chloride solution added; then without filtering barium hydroxide is added in slight excess, the precipitate filtered off, and washed. To the filtrate is added a little ammonium hydroxide, and ammonium carbonate solution until the barium is precipitated. This precipitate is filtered and washed, the filtrate evaporated to dryness, and carefully ignited below redness until all volatile matter is driven off. The residue is dissolved in a few cc. of water, and a few drops of ammonium carbonate solution added. The precipitate. if any, is removed by filtering and washing, and the filtrate evaporated in a small tared platinum dish, ignited below redness, and weighed. This gives the weight of the mixed chlorides. The residue is taken up with hot water, from 5 to 10 cc. of a 10% solution of platinic chloride added, and the whole evaporated to a sirupy consistency on the waterbath; it is then treated with 80% alcohol, the precipitate washed with 80% alcohol by decantation, transferred to a Gooch crucible, dried at 100° C., and weighed. The weight of the precipitate, multiplied by 0.10308, gives the weight of K₂O, and by 0.3056 the equivalent amount of KCl. The weight of KCl found is subtracted from the weight of the mixed chloride, the remainder being NaCl, which, multiplied by 0.5300 gives the weight of Na₂O in the sample.

^{*} U. S. Dept. of Agric., Div. of Chem., Bul. 13, part 5, p. 593.

Determination of Phosphoric Acid.—Method of the A. O. A. C.*— Mix 5 grams of the material with 10 cc. of magnesium nitrate solution. dry, ignite, and dissolve in hydrochloric acid. Take an aliquot part of the solution prepared above, corresponding to 0.25 gram, 0.50 gram, or I gram, neutralize with ammonia, and clear with a few drops of nitric acid. In case hydrochloric or sulphuric acid has been used as solvent, add about 15 grams of dry ammonium nitrate, or a solution containing that amount. To the hot solution add 50 cc. of molybdic solution! for every decigram of P.O. that is present. Digest at about 65° for an hour, filter. and wash with cold water, or preferably ammonium nitrate solution. Test the filtrate for phosphoric acid by renewed digestion and addition of more molybdic solution. Dissolve the precipitate on the filter with ammonia and hot water and wash into a beaker to a bulk of not more than 100 cc. Nearly neutralize with hydrochloric acid, cool, and add magnesia mixture from a burette; add slowly (about 1 drop per second), stirring vigorously. After fifteen minutes add 30 cc. of ammonia solution of density 0.96. Let stand for some time; two hours is usually enough. Filter, wash with 2.5% NH₃ until practically free from chlorides, ignite to whiteness or to a grayish white, and weigh.

Determination of Sulphuric Acid.—Provisional Method A. O. A. C. — Boil 5 grams of the powder gently for one and one-half hours with a mixture of 300 cc. of water and 15 cc. of concentrated hydrochloric acid. Dilute to 500 cc., draw off an aliquot portion of 100 cc., dilute considerably, precipitate with barium chloride, filter through a Gooch crucible, ignite, and weigh. Direct solution of the material without burning the organic matter was proposed by Crampton. The dextrose, formed by the action of the acid on the starch of baking-powders, does not interfere with the accuracy of the process.

Determination of Ammonia (present in the form of ammonia alum or ammonium carbonate). Mix 5 grams of the sample with 200 cc. of water, and add an excess of sodium hydroxide. Distill into standard acid, and determine the ammonia by titration.

^{*} U. S. Dept. of Agric., Div. of Chem., Bul. 46, p. 12; Bul. 107 (rev.), p. 4.

[†] Prepared as follows: Dissolve 80 grams calcined magnesia in nitric acid, avoiding an excess of acid, then add a little calcined magnesia in excess, boil, filter from the excess of magnesia, ferric oxide, etc., and dilute with water to 500 cc.

[‡] Reagent No. 53.

[§] Prepared by dissolving 100 grams of ammonium nitrate, Reagent No. 54, in 1 liter of water.

U. S. Dept. of Agric., Bur. of Chem., Bul. 65, p. 107; Bul. 107 (rev.), p. 178.

[¶] U. S. Dept. of Agric., Div. of Chem., Bul. 13, part 5, p. 596.

SEMOLINA, MACARONI, AND EDIBLE PASTES.

Semolina is the coarse meal ground from certain varieties of hard or "durum" wheats, grown originally in Italy, Sicily, and Russia, but at present in France and certain parts of the United States and Canada. This hard wheat is high in gluten, and especially adapted for the preparation of macaroni and the various pastes. A peculiar process is employed in preparing the wheat, whereby the husk is removed by wetting, heating, grinding, and sifting, the resulting meal or semolina, being in the form of small, round, glazed granules.

Italian Pastes.—Semolina furnishes the basis of the Italian edible pastes, being mixed with warm water, kneaded, and molded into various forms, either by pressure through holes in an iron plate, or otherwise, and finally dried. In parts of Italy juices of carrots, onions, and other vegetables are said to be mingled with the paste, but for local consumption only. Saffron is sometimes added to pastes for the purpose, so it is claimed, of imparting a spicy flavor, although the quantity used is often so small as to be apparent only to the eye, thus indicating that the real object of its addition is to impart a color in imitation of an egg paste.

Macaroni is the larger of the slender-tube or pipe-shaped products; vermicelli is the worm-shaped variety, produced when the holes in the plate are very small; spaghetti is the term applied to the cord-like paste intermediate in size between the others. A variety of Italian pastes or pâtés is made by rolling the kneaded semolina into thin sheets, and cutting out in shapes of animals, letters of the alphabet, etc.

The composition of some of these products is as follows:

-	No. of Samples.	Water.	Protein.	Fat.	Total Carbohy- drates.	Crude Fiber.	Ash.	Fuel Value per Pound. Cal's.
Semolina *	,	10.50	11.96	0.60	75-79	0.50	0.65	
Macaroni †	11	10.3	13.4	0.9	74.1		1.3	1665
Noodles †	2	10.7	11.7	1.0	75.6	0.4	1.0	1665
Spaghetti †	' 3	10.6	12.1	0.4	76.3	0.4	0.6	1660
Vermicelli †	15	11.0	10.9	2.0	72.0	••••	4.1	1625

^{*}Balland.

† Atwater and Bryant.

Noodles are a strap-shaped form of paste made in German house-holds as well as in factories. True, or egg-noodles, contain a certain percentage of eggs, while water-noodles are practically the same in composition as Italian pastes. The difference in composition between water-

noodles and noodles made with different numbers of eggs or egg yolks per German pound of flour, is shown by the analyses of Juckenack and Pasternack* given in the following table:†

of Eggs per of Flour.	Con	Composition of the Dry Matter.					Composition of the Dry Matter					
Number of Eg	Ash.	Total Phos- phoric Acid.	Lecithin Phos- phoric Acid.		Protein N×61	Number of Egg per Pound of	Ash.	Total Phos phoric Acid.	Lecithin Phos- phoric Acid.	Ether Extract	Protein N×61	
	%	%	%	0%	%		%	%	%	%	%	
٥	0.460	0.2300		0.66	12.00	0	0.460	0.2300	0.0225	0.66	12.03	
1	0.565	0.2716			12.99	1	0.488	0.2720	0.0518	1.57	12.37	
2	0.664	0.3110	0.0786	2.42	13.92		0.516	0.3127	0.0801	2.47	12.73	
3	0.758	0.3482	0.1044 *	3.24	14.81	3	0.542	0.3520	0.1 0 75	3.33	13.07	
12	1.426	0.6123	0.2875	7.94	21.09	12	0.745	0.6533	0.3171	8.64	15-71	

From these results it appears that the percentages of ash, total phosphoric acid, and protein are appreciably increased by the addition of each egg or egg yolk, while the percentages of lecithin-phosphoric acid and ether extract are more than doubled by the addition of the first egg, and are increased in corresponding proportion by the addition of two or more eggs.

The German Association of Food Chemists require that commercial egg-noodles contain at least 0.045% of lecithin-phosphoric acid, and 2.00% of ether extract, corresponding to the minimum in noodles with two eggs per half kilogram of flour.

Spaeth‡ considers that if the ether extract of noodles has an iodine number over 98, it is safe to assume that they contain no eggs or only traces.

In interpreting the results of analysis it should be remembered that fat may have been introduced in some form other than in eggs, and that the lecithin-phosphoric acid diminishes somewhat on long standing. Allowance should also be made for the variation in composition of the eggs and flour.

Of 22 brands of American noodles examined by Winton and Bailey only 5 appeared to be made with eggs; the lecithin-phosphoric acid in

^{*} Zeits. Unters. Nahr. Genuss., 3, 1900, p. 13; 8, 1904, p. 94.

[†] The German pound is approximately 468 grams; the avoirdupois pound is 454 grams.

[‡] Forsch. über Lebensm., 3, 1896, p. 49.

[§] Jour. Am. Chem. Soc. 1905, 37, p. 137; Rep. Conn. Exp. Sta., 1904, p. 138.

these ranged from 0.036 to 0.058, and the ether extract from 1.83 to 2.33 per cent, while in the other samples the lecithin-phosphoric ranged from 0.015 to 0.032 and the ether extract from 0.28 to 2.50 per cent.

Adulteration of Pastes.—Rice, corn, and potato flours have been used in the preparation of the cheaper varieties of semolina, but rarely in this country. A more common form of adulteration is the substitution of water-noodles for egg-noodles, artificial colors being used to carry out the deception. Substitutions of this kind are detected by determinations of lecithin-phosphoric acid and ether extract, supplemented by tests for artificial colors.

ANALYSIS OF PASTES.

Determination of Lecithin-phosphoric Acid.—Juckenack's Method.*
—Extract 30 grams of the finely ground material for 10 hours with absolute alcohol in a Soxhlet extractor at a temperature, inside the extractor, not below 55°-60° C. The extraction flask should be provided with a small quantity of pumice stone to prevent bumping during the boiling, and the extractor enclosed by asbestos paper, if the desired temperature is not readily maintained. After the extraction is completed, add 5 cc. of alcoholic solution of potash (prepared by dissolving 40 grams of phosphorus-free caustic potash in 1000 cc. alcohol), and distil off all the alcohol. Transfer the residue to a platinum dish by means of hot water, evaporate to dryness on a water bath, and char over asbestos. Treat the charred mass with dilute nitric acid, filter, and wash with water. Return the residue with the paper to the platinum dish, and burn to a white ash. Treat again with nitric acid, filter and wash, uniting the filtrates. Determine phosphoric acid by the usual method.

Detection of Artificial Colors in Pastes.—The following colors have been used in noodles and other pastes: turmeric, saffron, annatto, naphthol yellow (Martius yellow), naphthol yellow S, picric acid, aurantia, Victoria yellow, tartrazine, metanil yellow, azo yellow, gold yellow, and quinoline yellow. Of these naphthol yellow, picric acid, metanil yellow, and Victoria yellow are injurious to health, and their use is illegal in European countries as well as in the United States. Fortunately, they are rarely found in the products now on the market.

The detection of artificial colors is complicated by the presence of the natural coloring matter of the flour and the lutein of eggs. These are

^{*} Zeits. Unters. Nahr. Genuss., 3, 1900, p. 13.

conveniently extracted by ether, which does not remove the artificial colors, although most of them unmixed dissolve freely in this solvent.

Juckenack's Method.*—Thoroughly shake two portions of the finely ground material, each of about 10 grams, in test tubes with 15 cc. of ether and 15 cc. of 70% alcohol respectively, and allow to stand 12 hours.

- (a) If the ether remains uncolored or only slightly tinted and the material below it remains yellow, while the alcohol is distinctly colored and the material is decolorized, a foreign dye is indicated.
- (b) If both ether and alcohol are colored, either (1) lutein (egg color) alone, or (2) this with a foreign dye is present.
- 1. Treat a portion of the ether solution with dilute nitrous acid, according to Weyl. If the ether is not completely decolorized, a foreign dye is present.
- 2. If the deposit of material below the alcohol is decolorized, while that below the ether is colored, tests should be made for foreign dyes as follows: Shake the portion previously treated with ether with three or more fresh portions of the same solvent, until no more color is extracted, and then shake the residue with 70% alcohol and allow to stand 12 hours. After filtering, concentrate the solution slightly, acidify with hydrochloric acid, boil with sensitized wool, and identify the color in the usual manner (page 799).

Schlegel's Method.†—Extract 100 grams of the finely powdered material with ether in a continuous extraction apparatus, and shake the residue at frequent intervals for half a day with a mixture of 140 cc. of alcohol, 5 cc. of ammonia, and 105 cc. water. Filter, evaporate to remove alcohol and ammonia, acidify slightly with hydrochloric acid, and again filter. Boil the filtrate with sensitized wool, and identify the color on the dyed fiber by the usual tests (page 799).

Fresenius Method.‡—Extract 20 to 40 grams of the powdered material with ether in a continuous extraction apparatus. Dry the residue to remove ether, shake for 15 minutes with 120 cc. of 60% acetone, and allow to stand 12 to 24 hours. Filter, evaporate until the acetone is removed, and divide into two portions, a larger and a smaller. To the larger portion add sufficient acetic acid to dissolve flocks, and boil with sensitized wool. Remove natural coloring matter from the wool by boiling in dilute acetic acid. If after this treatment the wool is dyed.

^{*} Zeits. Unters. Nahr. Genuss., 3, 1900, p. 1.

[†] Untersuchungsanstalt, Nürnberg, Ber. 1906, p. 24.

[‡] Zeits. Unters. Nahr. Genuss., 13, 1907, p. 132.

the presence of a foreign color is indicated, which may be identified by the usual tests.

To the smaller portion of the aqueous solution, obtained after removal of the acetone as above described, add an equal volume of alcohol, heat to dissolve flocks, divide into four portions, and apply special tests to three of these, reserving the fourth for comparison. The natural color of the flour is decolorized by hydrochloric acid, intensified by ammonia, but not affected by stannous chloride, even on heating. Saffron reacts in a similar manner, but is only slightly bleached by the acid, and is not affected by the other two reagents.

Piutti and Bentivoglio Method.*—This method is especially designed to detect the four colors forbidden by Italian law, and to distinguish these from naphthol yellow S.

Boil 50 grams of the paste in 500 cc. of water, made alkaline with 2 cc. of concentrated ammonia water, add 60 to 70 cc. of alcohol, and continue the boiling 40 minutes. After filtering, acidify the liquid with 2 to 3 cc. of dilute hydrochloric acid and boil with 5 to 6 strands of sensitized wool, each strand weighing about 0.5 gram. Wash the wool, dissolve the color in dilute ammonia, and repeat the dyeing. After dissolving a second time in ammonia, evaporate the solution of the dye to dryness, avoiding as far as possible the formation of a skin, and take up the residue in water. If a skin has formed, filter and test the insoluble matter for metanil yellow with dilute hydrochloric acid, and for picric acid with ammonium sulphide.

To 1 cc. of the filtrate add stannous chloride solution and a little sodium hydroxide, or preferably sodium ethylate. If no red color forms, nitro-colors are absent; if, also, in another portion dilute hydrochloric acid produces no violet color, thus showing the absence of metanil yellow, no further test is necessary. In the presence of these colors, acidify the remainder of the solution with acetic acid, shake violently with carbon tetrachloride, and identify the color according to the following scheme:

- A. Color dissolves in carbon tetrachloride to colorless solution. Extract with very dilute ammonia, concentrate and divide into two parts.

^{*} Gaz. chim. Ital. 36, II, 1906, p. 385.

- B. Color is insoluble in carbon tetrachloride. Evaporate to dryness on water-bath, take up in water and divide into three parts.
 - I. Hydrochloric acid produces a violet coloration.... Metanil yellow,
 - 2. Ammonium sulphide produces a red brown coloration.

Picric acid.

3. Stir on a water-bath with zinc dust and ammonia, filter, treat with zinc dust and hydrochloric acid and again filter. (a) Potassium hydroxide produces a yellow coloration, and (b) ferric chloride an orange coloration.

Naphthol yellow S.

Schmitz-Dumont Test for Tropeolins.*—Moisten a small portion of the material with a few drops of dilute hydrochloric acid. The formation of a reddish or bluish color shows the presence of an azo color or some other coal-tar color.

Test for Turmeric.—Extract the color from the ground material by alcohol and identify by the boric acid test (page 780).

Shredded Wheat is a whole-wheat preparation, put out in the form of light biscuits built up of fine porous threads, not unlike those of vermicelli. The wheat, softened by boiling, is shredded by passing through a peculiar machine, after which the biscuits are made by lightly putting together the threads and by final baking. The comparative composition of shredded wheat and of typical whole wheat is thus shown by Wiley:†

Constituents.	Shredded Biscuit. Per Cent.	Typical Wheat. Per Cent.
Moisture Proteins Ether extract	10.57 12.06 1.03 2.65	10.60 12.25 1.75 1.75
Crude fiber. Carbohydrates other than fiber.	2.58 71.11	2.40 71.25

PREPARED CEREAL BREAKFAST FOODS.

The large number and variety of these preparations now on the market testify to the fact that the breakfast cereal forms a most important, as well as considerable, portion of our food supply. These foods are generally prepared from wheat, oats, and corn, and are, as a rule, remarkably pure and free from adulteration, though the food value of different varieties

^{*} Zeits. öffent. Chem., 8, 1902, p. 424.

[†] U. S. Dept. of Agric., Bur. of Chem., Bul. 13, p. 1337.

is often grossly misstated by their manufacturers. Formerly the break-fast food consisted entirely of the coarsely ground, generally decorticated, raw cereal grain, and required a long period of cooking to prepare it for use. At present many of the oat products, and to some extent also those of corn, rice, and wheat, are subjected to a more or less preliminary cooking and drying, whereby they are capable of being prepared for use in a much shorter time, and their keeping qualities are enhanced. The so-called rolled oats are prepared by softening the grains through steaming, after which they are crushed between rollers and afterwards dried. The steaming process is a typical one for various other cereals, though in some cases the heating consists in baking or kiln drying.

The effect of the preliminary cooking on the finished product varies somewhat according to whether dry or moist heat has been applied, and is chiefly noticeable in the altered character of the carbohydrates. In all cases the starch is rendered more soluble, whether by the conversion of a portion into dextrin and dextrose, or by a simple breaking down of the starch grains, as in the case of bread in baking.

In spite of the seemingly endless variety of the package cereals, they divide themselves as a matter of fact into a very few well-defined classes, the members of which differ but little from each other except in name.

First there are the raw cereal grains of the oat, wheat, and corn, prepared by simple crushing to various degrees of fineness, after decorticating; next comes the classes of partially cooked preparations of each of these grains, appearing in various forms of "flakes," "granules," "grits," etc., and again a class known as malted cereals, in which the moist, ground grain is mixed with malted barley, and, by controlling the temperature, a portion of the starch is converted to maltose and dextrin, after which the mixture is crushed between hot rollers and dried.

In the preparation of most of the corn breakfast products, such as samp and hominy, it is customary to remove the germ, which contains the oil and fat, lest the tendency of the latter to become rancid should result in the deterioration of the food. In wheat foods the germ is less often removed, and rarely, if ever, in oat preparations. The amount of fat found in the prepared cereal food as compared with that in the whole grain is of interest in this connection.

Composition of Some of the Common Breakfast Cereals.—The following analyses will serve to typify the various classes of these preparations as they appear on the market:

	Carbohydrates.									
	Water.	Fat.	Protein.	Total.	Soluble in Water.	Insoluble in Water	Crude Piber.	Ash.	Phosphorus.	Fuel Value per Gram. Calories.
WHEAT.*										
Wheatena	6.65			75.62		70.50	1.22	1.28	0.363	4343
Pettijohn's breakfast food				76.96	2.8	72.15	2.01	1.52	0.231	4174
Farina	10.94	1.56		75-91	3.2	72.12	0.59	0.69	0.153	4051
Cracked wheat		2.22	12.60	94-42	3-3	69.63	1.49	1.46	0.333	4236
Ralston's breakfast food	9.72	1.90	15.10	71.75	4.6	65.60	1.55	1.53	0.343	4158
Fould's wheat germ OATS.*	10.13	1.46	13.30	73-93	3.7	69.35	0.88	1.18	0.326	4087
Quaker	7-40	6.08	17.20	66.65	1.6	64.65	1.40	1.67	0.341	4673
Hornby's	7.63	7-35	17.82	65.47	1.3	62.74	1.43	1.73	0.443	4756
Buckeye	7-54	8.30	16.89	65.55		60.90				
Cerealine*	0.55	I . 2.1	0.00	78.75	7.1	70.93	0.72	0.56	0.102	4542
Velvet meal*	0.80	2.32				77-77				
Hecker's hominy †									3	0
Nichols' snow-white samp† MISCELLANFOUS.†				80.5			0.4			
Brittle bits	6.9	0.5	14.1	76.0			1.0	1.5	1]
Force		1.4	11.6	76.8			2.0	2.8		l
Grape-nuts	4.2						1.9	1.8	1	ŀ
Ralston's health barley food		1.0		75.8				1.0	1	i

^{*} Analyses made by Slosson, Wyoming Exp. Sta., Bul. 33.
† Analyses made by Merrill and Mansheld, Maine Exp. Sta., Bul. 84.

The methods of analysis employed for these preparations are the same as for ordinary cereals (p. 277), the sample being ground fine enough to pass through a 1-mm. sieve.

PREPARED FOODS FOR INFANTS AND INVALIDS.

In dealing with the composition and analysis of this class of proprietary foods more than ordinary care is necessary, in view of the fact that one or another of these preparations are frequently prescribed for the exclusive diet of those whose very life may depend on the character and suitability of the food to the case in hand. Many of these foods do, as a matter of fact, honestly fulfil the claims of their manufacturers, but others fall far short of so doing, so that it is hardly safe to use them unless some intelligent idea of their composition can be gained. It is not, as a rule, within the province of the analyst to furnish an opinion regarding the adaptability of a certain food to the requirements of an infant or invalid, but rather to provide the necessary data whereon such an opinion may be intelligently based.

A simple statement of moisture, fat, protein, carbohydrates (by difference), and ash, which in the case of ordinary foods would often be sufficient, would be obviously inadequate in expressing the analysis of an infant food, since it is of much more vital importance than in other foods to know the solubility of the food itself, and, to as great an extent as possible, the character of the carbohydrates.

The chief ingredients of many of these preparations are wheat, or mixed cereals high in starch. Many of the foods are, according to the directions, to be used practically without cooking, but by simply mixing with milk or water, and, in some cases, bringing to the boiling-point. Hence the degree of conversion which the raw starch has undergone in the process of manufacture of the food should, if possible, be ascertained as a prime factor in judging of its character and adaptability to the needs of the young child and of the sick. Incidentally it should be said that few if any of the infant foods, even those whose high character has long been established by continued trial, conform very closely to the composition of woman's milk, which was long accepted as the true standard on which to base their efficiency. Hence it is no easy task to pass judgment on a particular food from its chemical composition alone without trial. nor is it right to unqualifiedly condemn in all cases food high in insoluble carbohydrates, since there are undoubtedly many instances in which such foods are successfully used.

Classification and Preparation of Infants' Foods.—These foods may for convenience be divided into two main classes, viz., farinaceous foods, or those which are prepared wholly or chiefly from one or more cereal grains, and lactated foods, or those in which cow's milk forms the basis, but which may contain in addition thereto various other substances, such as cereals, sugars, etc.

The farinaceous foods, which are usually directed to be mixed with milk before using, may be further subdivided into (a) those that consist chiefly of unconverted starch, (b) those whose starch has been nearly all hydrolyzed to soluble form in the process of manufacture, and (c) those which contain much unconverted starch, but in addition thereto diastase or some other ferment, which, when the food is mixed with warm water or milk, is supposed to convert all the starch to soluble form.

The unconverted starch foods are nearly all made up of baked dry flour, chiefly that of wheat, but sometimes a mixture of cereals (as oats, barley, and wheat) and sometimes oats or barley alone. The baking breaks down to some extent the starch grains, as in the case of bread or crackers, but does not actually convert much of it to sugar.

The soluble farinaceous foods are usually prepared somewhat as follows: A mixture of ground wheat and barley malt (with sometimes a little wheat bran) is mixed with water to form a paste, and a little bicarbonate of potash added. The mixture is heated at 65° C. for sufficient time to convert the starch, after which it is exhausted with warm water, the extract being strained, and the filtrate evaporated to dryness to form the food. The sugars of such foods consist largely of maltose mixed with dextrin.

The farinaceous foods, which depend for the conversion of their starch on the method of cooking or heating before serving, are usually mixtures of wheat or other cereal flour with malt or pancreatic extract.

The milk foods are variously prepared, either by the simple desiccation of cow's milk (usually previously skimmed) or, when whole milk is used, by mingling the desiccated milk with sugars or baked cereal flour. Sometimes desiccated milk is used in mixture with a dried extract of malted cereals. In fact all sorts of mixtures are found on the market, involving, however, in nearly all cases, one modification or another of the above general processes of preparation.

Composition.—Few complete analyses of these classes of foods have recently been made. Among the best are those of McGill,* from whose work the following figures have been selected, illustrating typical examples of foods on the market:

	Number of Analyses.	Moisture.	Fat by Petro- leum Ether.	Loss to Alcohol.	Loss to Water.	Sum of Losses to Alcohol and Water.	Albuminoids, N×6.25.	Ash.
Farinaceous foods: Imperial granum. Ridge's food. Mother's food. Robinson's barley. Mixed foods:	5 9 2 7	6.04 8.12 9.99 9.41	0.72 0.48 0.13 0.41	0.34		3.94 5.02 8.83 2.91	13.77 13.83 8.60 7.46	0.53
Horlick's malted milk. Lactated food. Mellin's food. Nestlé's milk food. Reid & Carnrick's baby food.	9 12 8 9 2	2.55 5.77 4.72 2.18 5.69	1.41 0.48 0.30 4.45 2.18	28.24 39-54		63.87 32.90 82.0 43.84 38.21	10.01	2.57 3.50 1.60

^{*} Canadian Dept. of Inland Rev., Bul. 59.

	Starch, Fiber. etc., by Differ- ence.	Maltose.	Lactose.	Cane Sugar.	Remarks.
Farinaceous foods:				i	
Imperial granum	76.60				Wheat starch
Ridge's food.	72.01		i	l	Wheat starch
Mother's food	69.24	 		3.00	Corn and wheat starch
Robinson's barley	78.66				Barley starch
Mixed foods:	1		Ì		
Horlick's malted milk	15.68	49.00		8.00	
Lactated food	47.72		30.00	Trace	Wheat starch
Mellin's food		50 to 60			·
Nestlé's milk food	35 - 34			36.34	Wheat starch
Reid & Carnrick's baby food	34-54		30.00		

Diabetic Foods..—Gluten flour and similar preparations are primarily intended for the use of diabetics, from whose dietary carbohydrates must be excluded.

The following analyses of commercial gluten preparations were made by Woods and Merrill.*

	Protein.	Fat.	Carbohy- drates.	Ash.
"Cooked gluten". Whole-wheat gluten "Glutine" Breakfast cereal gluten. Plain gluten flour Self-raising flour	17.89 15.31 43.70	3.86 5.20 0.99 1.60 1.20	76.80 73.85 82.52 44.40 34.50 53.20	2.46 3.06 1.17 0.70 0.60 3.80

Many brands of gluten flour are put on the market by dealers in so-called "health foods," and in many cases are represented to be practically free from starch. Thirteen samples of gluten flour were analyzed by the author in 1899, varying in price from 11 to 50 cents per pound. Of these, 3, the product of one manufacturer, contained less than 1% of starch, 3 contained from 10 to 20 per cent, while 7 contained from 56 to 70 per cent of starch, the substance which, of all others, the diabetic patient tries to avoid. Some of these preparations were little better than wholewheat flour. An analysis of one of them, known as "Pure Vegetable Gluten," and sold for 50 cents per pound, and of two similar diabetic flours reported by Winton follow:

^{*} Maine Exp. Sta. Buls. 55 and 75.

	"Pure Vegetable Gluten."	" Diabetic Food."	" Diabetic Flour."
Moisture	10.78	12.67	9.26
Ash	2.20	0.43	1.30
Fat	3.25	0.90	2.21
Protein	14.25	11.37	14.25
Crude Fiber	•••	0.25	1.03
Sugars	1.70 }		
Dextrin	2.55 }	71.51	66.63
Starch	56.55		
Undetermined	8.72	2.87	5.32
	100.00	100.00	100.00

Winton has reported the following analyses of flours and meals well suited for the preparation of diabetic biscuit, and of the biscuit made from two of these by a cook in the family of a diabetic patient:

	Moisture.	Ash.	Protein, (N×64).	Crude Fiber.	Nitrogen-free Extract.	Fat.	Starch, Sugar, and Dextrin.
Gluten nour Calc. wat	al 10.12 er-free	0.24	85.38 95.00		3.69 4.11	0.62	4.46 4.96
	al 25.58 er-free		50.91 68.41		3.18 4.27		
	al 7.75 er-free		39.87	3.85	25.09 27.20	19.06	
	al 27.66 er-free		16.71 23.10		12.84	35.91	
	er-free10.01	2.46	85.56			0.50 0.56	
Almond meal In origina	er-free8.51	6.42	50.62 55-32	2.86	15.96 17.45		7.18

In the analysis of diabetic foods, the determination of starch, sugar and dextrin together is of greater value than of starch alone, since all three classes of carbohydrates are about equally injurious to diabetics, the starch and dextrins being converted into sugars by the digestive fluids. The nitrogen-free extract of cereal preparations corresponds closely with the sum of the starch, sugar and dextrin, but in the case of soja bean meal, almond meal and other products of legumes and oil seeds, as well as vegetables, it is considerably greater, as it includes pentosans and other substances.

METHODS OF ANALYSIS.

The sample is prepared for analysis by grinding it sufficiently fine in a mortar or mill to pass through a 1-mm. sieve. Moisture, fat, ash, and nitrogen are determined as in the regular methods for cereals (pp. 277, 278).

In determining loss of weight due to solubility of the sample in alcohol and water, proceed as follows:* The fat-free residue left in the Soxhlet apparatus, after extraction with ether or petroleum ether, is subjected to further extraction with 05% alcohol, till all soluble matter has been extracted. If 5 grams of the sample were originally taken for the fat extraction, this operation would require about five hours. Evaporate the alcoholic extract to dryness, and weigh the residue as in the case of the ether extract. Dry the residue left in the Soxhlet from the alcoholic extract, or a portion thereof, in a platinum dish over the water-bath, cool, and weigh. Transfer to a Gooch crucible, provided with asbestos and previously tared, a portion, the relation of which to the original weight taken is calculated from the moisture, ether, and alcohol extracts as previously determined. Pass through the contents in the Gooch by suction from 200 to 300 cc. of cold water at room temperature, dry the Gooch and its contents at 100° to constant weight, cool and weigh, thus determining the solubility of the sample in water.

According to McGill, five hours' extraction with alcohol under the above conditions removes all cane sugar, but probably not all the lactose, maltose, and dextrose, if a considerable quantity of these sugars is present. Water dissolves the dextrin and gum and such of the sugar as escapes solution in the alcohol, hence the sum of the alcohol and water extract is of value. In the calculation of the starch, fiber, etc., by difference, it should be borne in mind that the result is only approximate, by reason of the fact that the small amount of soluble albuminoids (which McGill states never exceeds $2\frac{1}{2}\%$) are reckoned in, hence a small error is introduced, which could be corrected, if considered worth while, by determining the amount of soluble albuminoids.

Separation of the Carbohydrates can be effected by Stone's method (pp. 295, 296), but a very satisfactory idea of the solubility of these foods, which is of chief importance, can be gained by the much simpler modified method of McGill, as described in the preceding paragraphs.

^{*} McGill, Canada Inland Rev. Dept., Bull. 58.

Starch, Sugar, and Dextrin are determined together in diabetic preparations by the diastase method (p. 283) omitting the preliminary washing with dilute alcohol.

Cold-water Extract.—The equivalent of 10 grams of the moisture-free substance, finely ground, is weighed in a tared flask, and water added in several portions with gentle shaking till the contents of the flask weigh 110 grams. The flask is then corked and vigorously shaken at intervals during six or eight hours and allowed to stand over night. The supernatant liquid is then decanted into the large tubes of a centrifuge, and whirled till the sediment settles out. The comparatively clear liquid may then be readily filtered. 20 cc. of the filtrate, corresponding to 2 grams of the original sample, are then transferred to a tared dish, evaporated to dryness, and dried to constant weight, as in the determination of the total solids.

Additional information may be gained from the specific gravity of the 10% solution of the cold-water extract, best obtained by means of a pycnometer.

Reducing Sugars are determined in an aliquot part of the above 10% solution, diluted to proper strength.

Effects of Subsequent Heating.—It is hardly fair in the case of those farinaceous foods which, according to directions, are to be subsequently subjected to heating, or boiling with water or milk, to condemn them as containing much insoluble matter, without comparing the figures expressing results of the analyses of the raw foods, calculated to the water-free basis, with those obtained on analyzing the food after boiling or otherwise cooking with pure distilled water, for a length of time specified in the directions, and afterwards drying. It is possible that the presence in the food of diastase, or other ferment, may be depended on to hydrolyze a whole or a portion of the starch, and only by such comparison will this be shown.

Microscopical Examination of the food is of value in determining its general character, showing especially whether or not starch is present in its original form, or has been converted in whole or in part. The particular varieties of cereal grain employed are generally evident, as well as the presence and proportion of the different tissues of the grain.

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CHAPTER XI.

TEA, COFFEE, AND COCOA.

TEA.

Nature and Classification.—Tea consists of the prepared leaves or leaf buds of Camellia Thea also known as Thea chinensis.

The best teas are made from young leaves only, the Chinese teas being classified with reference to the age and position of the leaf on the young shoot. Thus, the very choicest Chinese tea, rarely found outside of China, is prepared from the youngest or end leaves of the shoot, which are scarcely more than buds, and form the tea known as pekoe tip, or flowery pekoe. The next leaves are the orange pekoe and pekoe, which produce a very high grade of tea, while next in order as to age, size, and grade of leaf are the souchong 1st and 2d, and the congou, producing teas called by the same names.

More than 50% of the tea consumed in the United States comes from China, and over 40% from Japan, the remainder being derived largely from India, Ceylon, and other East Indian ports.

In the manufacture of tea the fresh leaves, which are nearly 80% water, are rolled, withered by exposure to light, heat, and air, and finally dried or "fired" by treatment with artificial heat over charcoal fires, or in properly constructed furnaces.

Teas are divided into two groups, black and green, which differ from each other, not as formerly supposed in being derived from different plants, but in their process of manufacture, the same species of plant furnishing both varieties. Genuine green tea is prepared by first steaming and afterwards drying the leaves while still fresh, thus retaining the bright color. Black tea is allowed to undergo oxidation or fermentation by exposure to the sun, which gradually turns the leaves black. Less tannin is present in black tea than in green.

Composition of Tea.—König gives the following composition of fully developed tea leaves, being the mean of 50 to 70 analyses:

Water.	Nitroge- nous Sub- stances.	Theine.	Essential Oil.	Pat, Chlo- rophyl, and Wax.	Gum, Dextrin, etc.	Tannin.	Pectin, etc.	Crude Piber.	Ash.
9-51	24.50	3.58	0.68	6.39	6.44	15-65	16.02	11.58	5.65

Though the nitrogenous substances of tea predominate in amount over any other class of constituents, yet, with the exception of theine or





Fig. 72.—a, Flowery Pekoe; b, Orange Pekoe; c, Pekoe; d, Souchong, 1st; c, Souchong, 2d; f, Congou; a, b (when mixed together), Pekoe; a, b, c, d, c (when mixed together), Pekoe Souchong. If there be another leaf below f, it is termed Bohea. At base of leaves are buds 1, 2, 3, 4, from which new shoots spring. (After Money.)

caffeine, they have been little studied. Theine, tannin, and essential oil give to the infusion of tea its chief characteristics.

Zollinski * gives the following summarized results of analyses of a number of the cheaper grades of Chinese black tea:

^{*} Zeits. anal. Chem., 1898, 37, 365.

	Water.	Total Nitrogen.	Albumin- oid and Amido- nitrogen.	Protein, N×6.25.	Theine.	Ash.	Soluble Ash.	Insoluble Ash.
Maximum Minimum Average	11.57	4.12	3.78	23.83	2.06	6.78	31.17	61.03
	9.96	3.76	3.37	21.06	1.14	4-79	28.13	57.74
	10.58	3.93	3.52	22.01	1.55	5-94	29.67	59.75

A very complete series of analyses of tea was made by Joseph F. Geissler in 1884,* from which the following summaries are taken:

		Number of Analyses.	Moisture.	Half-hour Extract.	Total Extract.	Insoluble Leaf.	Tannin.	Theine.	Total Ash.	Soluble Ash.	Insoluble Ash.	Ash Insol. in Acid.
Indian:	Maximum	6	6.19	39.66 37.80	45.64 41.32	53.07 48.53	18.86 13.04	3-3 1.8	5·79 5·42	3.68	2.22 1.93	.296
	Average		5.81	38.77	42.94	51.24	14.87	2.7	5.62	3-52	2.12	. 178
Oolong:	Maximum	13	6.88	44.02	48.87	53-15	20.07	3.50		3-71	3.17	.838
	Minimum	••	5.00	34.10	40.6	44.8	11.93		5-44	2.60	1.84	. 266
_	Average	••	5.89	37.88	43.32	50.7	16.38	2.32	5.81		2.68	. 507
Congou:	Maximum	11					13.89	2.87		3.52	3.86	1.31
	Minimum		7.65	23.48	27.48	54 - 5	8.44	1.70	5.52	2.28	1.90	. 32
	Average	••	8.37	28.40	34 - 35	57.2	11.54	2.37	5-75	3.06	2.68	-425

Kenrick † gives the following averages of a series of analyses of tea made by him in 1891:

		Substances Extracted by 10 Minutes' Infusion.					Ash.		Total o Tannin.
	Number of Analyses.	Total Solids.	Tannin.	Theine.	Moisture.	Soluble Ash.	Insoluble A	Total Ash.	Ratio of To Solids to T
Congou tea	10 3 2 13 18 2 5	23.37 38.53 27.45 23.76 30.07 28.55 34.22	7.85 5.40 9.38 8.00	2.98 2.68 2.82 2.45 2.30	7.60 5.75 6.31 6.54 4.00 4.72 5.40	3.55 3.69 3.34 3.53 3.62 3.36 3.83	2.28 2.16 1.88 2.37 2.73 3.70 2.10	5.83 5.84 5.22 5.90 6.35 7.06 5.93	4.51 3.81 3.50 4.40 3.20 3.57 3.12

The ash of many genuine teas has been examined by Battershal‡ with the following results:

^{*} Am. Grocer, Oct. 23, 1884.

[†] Canada Inland Rev. Dep. Bul. 24.

[‡] Food Adulteration and its Detection.

	Oolong. Average of 50 Samples.	Japan,	Spent Black Tea.
Total ash	6.04	5-55	2.52
Soluble in water	3-44 57.∞	3.60 64.55	0.28
COMPOSITION	ON.		ı
Silica	11.30	9.30	27-75
Chlorine	1.53	1.60	0.79
Potash	37.46	41.63	į
Soda	1.40	1.12	1.
Ferric oxide	1.80	1.12	16.00
Alumina	0 0 1	4.26	1, 10.00
Manganic oxide	2.10	1.30	
Lime	9-43	8.18	19.66
Magnesia	8.00	5-33	11.20
Phosphoric acid	12.27	16.62	15.80
Sulphuric acid	4.18	3.64	1.10
Carbonic acid	5.40	5-90	6.70
	100.00	100.00	99.00

Kozai* gives the following as the results of analyses made by him of Japanese teas:

	Unprepared Leaves.	Green Tea.	Black Tea.
Caffeine or theine	3.30	3.20	3.30
Ether extract	6.49	5.52	5.82
Hot-water extract	50.97	53-74	47-23
Tannin (as gallotannic acid)	12.91	10.64	4.89
Other nitrogen-free extract	27.86	31-43	35 - 39
Crude protein	37 - 33	37 - 43	38.90
Crude fiber	10.44	10.06	10.07
Ash	4-97	4.92	4-93
Albuminoid nitrogen	4.11	3-94	4.11
Caffeine nitrogen.	0.96	0.93	0.96
Amido-nitrogen	0.91	1.13	1.16
Total nitrogen	5-97	5.99	6.22

PROXIMATE COMPONENTS AND ANALYTICAL METHODS.

Preparation of Sample.—Grind the material so as to pass a sieve with holes 0.5 mm. in diameter.

Moisture, Ether Extract, and Crude Fiber are determined in the same weighed portion of 2 grams, following the methods described under cereals (p. 277).

^{*} Bul. 7, Imperial College of Agriculture, Japan.

Protein.—Determine total nitrogen by the Kjeldahl or Gunning method; from this subtract the nitrogen due to caffein (obtain by dividing by 3.464) and multiply the difference by 6.25.

Total Ash.—Burn 2 grams of the material to a white ash in a platinum dish at a faint red heat. The total ash of pure tea should not be less than 4 nor more than 7%.

Soluble and Insoluble Ash.*—The total ash, as obtained above, is transferred to a beaker with hot water and boiled for some time, after which it is poured upon a filter and the residue washed with hot water. The residue is then dried, ignited at a faint red heat in a platinum dish, cooled, and weighed, thus giving the amount of insoluble ash. The soluble ash is calculated by difference from the total and insoluble ash.

Ash Insoluble in Acid.*—The portion of the ash insoluble in water is washed upon a tared filter with hot 10% hydrochloric acid and further washed with the latter reagent till the acid-soluble matter is dissolved out, after which it is washed with water, dried, and weighed.

Alkalinity of Ash.*—This is expressed in terms of cc. of tenthnormal acid required for the ash of 1 gram of sample.

Soluble Ash.—Cool the filtrate from the determination of insoluble ash, as described above, and titrate with tenth-normal hydrochloric acid, using methyl orange as an indicator.

Insoluble Ash.—Add excess of tenth-normal hydrochloric acid (usually 10 to 15 cc.) to the ignited insoluble ash as obtained above in the platinum dish, heat to the point of boiling over an asbestos plate, cool, and titrate excess of hydrochloric acid with tenth-normal sodium hydroxide, using methyl orange as an indicator.

Essential Oils.—Distil 100 grams of the tea with 800 cc. of water, and shake out the distillate with several portions of ether. The residue from the combined ether extracts contains the volatile oil.

Insoluble Leaf.†—Boil 2 grams of the tea in a 500-cc. Erlenmeyer flask over a low flame with 200 cc. of water, replacing from time to time by addition of hot water the loss from evaporation. Filter through a tared filter, and wash the residue until the filtrate measures 500 cc., stirring the contents of the filter throughout the process to facilitate filtering. Reserve filtrate for determination of tannin and theine. Dry the filter and residue until dry to the touch, transfer to the weighing bottle, and dry to constant weight at 100° C. If the amount of insoluble

^{*}A. O. A. C. Method, U. S. Dept. of Agric., Bur. of Chem., Bul. 107 (rev.), p. 69. † Doolittle and Woodruff, A. O. A. C. Proc. 1906, U. S. Dept. of Agric., Bur. of Chem., Bul. 105, p. 48. Winton, Ogden and Mitchell, Conn. Exp. Sta. An. Rep., 1898, p. 132.

leaf is above 60%, the presence of spent or exhausted leaves may be suspected.

Extract.—By this term is meant the total amount of water-soluble matter in tea, including such compounds as tannin, caffeine, albuminous matter, dextrin, gum, certain parts of the ash, etc.

The value of a tea from a food standpoint depends obviously upon the character and amount of the extract, rather than on the composition of the dry tea. The relative composition of the extract and of the insoluble leaf, as found by Eder, is given in the following table.

	Extract.	Insoluble Leaf.
.	Per Cent.	Per Cent.
Dry matter	40.	60.
Nitrogenous substances	12.	12.7
Theine	2.	1
Tea oil	0.6	
Resin, chlorophyll, etc		7.2
Tannin	10.	1
Extractives	12.	10.
Ash	1.7	2.3
Potash	0.94	0.29
Lime	0.04	0.58
Phosphoric anhydride	0.13	1.03
Silica.	0.21	0.68

Determination.—The sum of the percentages of insoluble leaf and moisture subtracted from 100 gives the percentage of extract.

Tannin.—Proctor's Modification of Lowenthal's Method.*

- Reagents: (1) Potassium permanganate solution containing about 1.33 grams per liter.
 - (2) Tenth-normal oxalic acid solution (6.3 grams per liter).
 - (3) Indigo carmine solution, containing 6 grams indigo carmine (free from indigo blue) and 50 cc. concentrated sulphuric acid per liter.
 - (4) Gelatin solution, prepared by soaking 25 grams gelatin for an hour in a saturated sodium chloride solution, heating till the gelatin is dissolved, and making up to a liter after cooling.
 - (5) Mixture of 975 cc. saturated sodium chloride solution and 25 cc. concentrated sulphuric acid.
 - (6) Powdered kaolin.

Obtain the value of the potassium permanganate solution in terms of the tenth-normal oxalic acid solution.

^{*} U. S. Dept. of Agric., Div. of Chem., Bul. 13, part 7, p. 890; Bul. 107 (rev.), p. 150.

Boil 5 grams of the powdered tea for half an hour with 400 cc. of water, cool, transfer to a graduated 500-cc. flask, and make up to the mark. To 10 cc. of the infusion (filtered if not clear) add 25 cc. of the indigo carmine solution and about 750 cc. of water. Then add from a burette the potassium permanganate solution, a little at a time while stirring, till the color becomes light green, then cautiously drop by drop till the color changes to bright yellow,* or further to a faint pink at the rim. The volume in cubic centimeters of permanganate furnishes value a of the formula.

Mix 100 cc. of the clear infusion of tea with 50 cc. of gelatin solution, 100 cc. of salt acid solution, and 10 grams of kaolin, and shake several minutes in a corked flask. After settling, decant first the clear supernatant liquid through a filter, and finally bring the precipitate upon the filter. Mix 25 cc. of the filtrate (corresponding to 10 cc. of the original infusion) with 25 cc. of the indigo carmine solution, and about 750 cc. of water, and titrate with permanganate as before. The volume in cubic centimeters of permanganate used gives value b.

a=quantity of permanganate solution required to oxidize all oxidizable substances present.

b=quantity of permanganate solution required to oxidize substances other than tannin.

 $\therefore a-b=c$, permanganate required for the tannin. Assuming that 0.04157 gram tannin (gallotannic acid) is equivalent to 0.063 gram oxalic acid, the tannin in the tea is readily calculated.

As recommended by Doolittle and Woodruff† the determination may be conveniently made on aliquot portions of the solution obtained in the determination of insoluble leaf.

Method of Fletcher and Allen.‡—This method depends upon the precipitation of the tannin and other astringent matters in tea infusion by lead acetate, the point of complete precipitation being indicated by an ammoniacal solution of potassium ferricyanide.

Five grains of neutral lead acetate are dissolved in water, made up to 1 liter, and after standing the solution is filtered.

As an indicator, 0.05 gram of pure potassium ferricyanide is dissolved in 50 cc. of water, and an equal volume of concentrated ammonia-

^{*} Various shades of color may be produced, but the same shade should obviously be adopted as an end-point by the operator as when standardizing.

[†] A. O. A. C. Proc. 1906, U. S. Dept. of Agric., Bur. of Chem., Bul. 105, p. 49.

[‡] Chem. News, XXIX, 169, 189.

water is added. This indicator produces a red coloration with tannin, gallic acid, or gallotannic acid in solution, being so sensitive that a drop of the indicator will detect 1 part of tannin in 10,000 parts of water.

Three separate quantities of 10 cc. each of the standard lead acetate solution, as above prepared, are measured into as many beakers, and each diluted to 100 cc. with boiling water. Two grams of powdered tea are boiled in 250 cc. of water, and varying quantities of this decoction are measured from a burette or pipette into the beakers containing the lead solution, the first beaker receiving, say, 12 cc., the second 15 cc., and the third 18 cc., in the case of black tea, and, with green tea, 8, 10, and 12 cc., respectively.

About 1 cc. of each of these trial quantities is removed from the various beakers by means of a pipette, passed through small filters, and tested with the ammoniacal ferricyanide indicator, the drops of filtered solution being allowed to fall directly on spots of the indicator, previously placed on a white slab or plate.

It is thus easy to ascertain the *approximate* amount of tea solution which it is necessary to add to produce a pink coloration with the indicator, so that by repeated tests, nearly the right amount may be added at once. If no coloration in a given case is produced when a drop of the filtrate from the solution in the beaker is allowed to fall on the drop of indicator solution, a little more of the tea decoction is added, and this process is repeated until the pink color is apparent.

It should be noted how much of the tea decoction is necessary to add to 100 cc. of pure water, that a drop of the solution may produce the pink coloration with the ferricyanide, and this amount should be subtracted from the amount of decoction found necessary to add to the known lead solution in the beaker. It was found by repeated experiment that 10 cc. of lead solution would precipitate 0.01 gram of pure gallotannic acid; hence, carrying out the process exactly as above described, 125 divided by the number of cubic centimeters of tea decoction required gives the percentage of tannin in the sample.*

Theine or Caffeine (C₈H₁₀N₄O₂).—This alkaloid when pure exists in white silky needles. It is odorless and sparingly soluble in cold water,

^{*} This process estimates the total astringent matter, all of which is counted in as tannin.

but more so in hot. It is less soluble in alcohol, and almost insoluble in ether. It readily dissolves in chloroform. It is present in tea, coffee, and kola. Graf* has shown that the amount of caffeine present in tea is in most cases proportional to the commercial value and quality.

Detection.—Caffeine may be detected, if present in a suspected residue, by the so-called "murexid test," which is made with the material in a solid state, or with the residue from the evaporation of a liquid. A small quantity of the solid or powdered material is heated in a white porcelain dish and covered with a few drops of strong hydrochloric acid, after which a fragment of potassium chlorate is immediately added. The mixture is then evaporated to complete dryness on the water-bath, whereupon, if caffeine is present, a reddish-yellow or pink color is produced. After cooling, the residue is treated with a very little ammonia water applied on the point of a stirring-rod. In the presence of caffeine, a purple color (that of murexoin) is produced on application of the ammonia.

Determination of Theine or Caffeine.—Dvorkovitsch Method.†—Digest 10 grams of the powdered tea with 200 cc. of boiling water for 5 minutes and decant the solution; repeat the treatment twice, and boil the residue with 200 cc. of water. Make up the combined solutions to 1000 cc. and extract a portion with petroleum ether to remove fat, etc. To 600 cc. of the fat-free solution (equivalent to 6 grams of tea) add 100 cc. of 4% barium hydroxide solution, mix and filter. To 583 cc. of the filtrate (equivalent to 5 grams of tea) add 100 cc. of a 20% solution of sodium chloride, and extract three times with chloroform. Distil the greater part of the chloroform from the combined extracts, place the residue in a tared dish, evaporate the remainder of the chloroform, dry at 100° C., and weigh. The caffeine is usually of sufficient purity to render a nitrogen determination unnecessary.

Doolittle and Woodruff[‡] proceed as follows: Extract in a separating funnel with petroleum ether 225 cc. of the filtrate from the determination of insoluble leaf (p. 369) made up to 500 cc. To the fat-free portion add 50 cc. of a 4% barium hydroxide solution, shake well,

^{*} Forsch, Ber., 4, 1897, pp. 88, 89.

[†] Ber. d. chem. Ges., 24, 1891, p. 1945; U. S. Dept. of Agric., Bur. of Chem., Bul. 107 (rev.), p. 150.

[‡] loc. cit.

and filter. To the filtrate add 50 cc. of a 20% sodium chloride solution and proceed as above described.

Modification of Stahlschmidt's Method.—Six grams of finely powdered tea are boiled in a flask with several successive portions of water for ten minutes each, and the combined aqueous extracts thus obtained are made up to 600 cc. with water. Four grams of powdered lead acetate are added to the decoction, which is then boiled for ten minutes, using a reflux condenser, or making up the loss by occasional addition of water. The solution is then poured upon a dry filter, and 500 cc. of the filtrate, corresponding to 5 grams of the tea, are evaporated to about 50 cc. and enough sodium phosphate added to precipitate the remaining lead. The solution is then filtered, and the precipitate thoroughly washed, the filtrate and washings being evaporated to about 40 cc. Finally, the solution thus concentrated is extracted with chloroform in a separatory funnel for at least four times, and the chloroform extract evaporated to dryness, leaving the caffeine, which is dried to constant weight at 75° and weighed.

ADULTERATION OF TEA.

Facing.—The most common form of tea adulteration, if such it may be called, is the practice of "facing" the dried leaves, or treating them with certain pigments and coloring materials to impart a bright color or gloss to the tea, thus causing an inferior grade to appear of better quality than it really is. This practice is more often applied to green tea. The materials for facing include such substances as Prussian blue, indigo, plumbago, and turmeric, often accompanied by such minerals as soapstone, gypsum, etc. Only a small amount of foreign material is actually added to the tea, but the adulteration consists in the deceptive appearance imparted thereto.

Battershal has examined various samples of the preparations used in Japan for facing tea. He found in one case the following composition: Soapstone, 47.5%; gypsum, 47.5%; Prussian blue, 5%. Another sample consisted of soapstone, 75%; indigo, 25%. A third was composed of soapstone, 60%, and indigo, 40%. In applying the facing to the tea, the latter is first heated in an iron pan over the fire, the facing

mixture is then added while still hot, and the whole is stirred briskly till the desired color is imparted. The Chinese and Japanese do not face the tea which they themselves consume, but only that intended for export trade.

The microscope furnishes the most ready means of detecting turmeric and plumbago. The latter is detected by the bright glossy particles, evident when a thin section of the tea leaf is examined under the microscope.

Prussian blue and indigo are also evident by the microscopical appearance of the particles, detached by shaking the leaves in water. Prussian blue or ferric ferrocyanide is detected by the transparent bright-blue particles, while indigo, when viewed under the microscope, is more of a greenish blue. The detached particles of coloring matter often rise to the surface of the liquid, when the leaves are shaken in hot water, and for microscopical examination may be floated upon a glass slide. The color of Prussian blue is discharged by treatment with sodium hydroxide, while that of indigo is not. Prussian blue, if present, may be chemically detected in the sediment as above obtained, by dissolving in hot alkali, acidifying with hydrochloric acid, and then adding a drop of ferric chloride. A blue precipitate is indicative of the ferric ferrocyanide.

Such minerals as gypsum and soapstone are readily separated as a sediment by shaking the leaves in water, and the sediment is examined by the appropriate qualitative methods for these substances.

Spent or Exhausted Leaves.—These consist of leaves of tea that have been previously steeped or infused, and afterwards rerolled and dried. Such leaves are sometimes mixed with tea as an adulterant. Any considerable admixture of spent leaves is evident, both by the extremely low ash, and the abnormally small proportion of water-soluble ash in the sample. It is rare that the total ash of genuine tea is under 5%, while the soluble ash is seldom less than 3%.

The ash of spent tea leaves sometimes runs as low as 2.5%, of which generally not more than 0.3 to 0.8 per cent is soluble. Spent leaves are also naturally low in tannin and in extract.

If the extract is much below 32%, spent leaves may be suspected. Allen has suggested the following formula for determining the percentage of spent leaves, E, in a sample of tea, R being the percentage of extract:

$$E = \frac{(32 - R)100}{30}$$

The use of spent or exhausted leaves as an adulterant is very rare at present, though formerly of common occurrence.

Foreign Leaves as a Substitute for Tea.—This sophistication is not common, but the detection of leaves other than tea is readily accomplished by a careful examination of the shape and character of the leaves. For this purpose the dried leaves are opened out by soaking a short time in hot water, after which they are spread upon a glass plate, and examined by the aid of a magnifying-glass.

The genuine tea leaf (Fig. 73) is very characteristic, and is readily distinguished from other leaves. It is oval or lanceolate, 5 to 8 cm. long

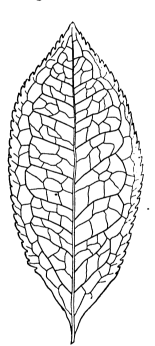


Fig. 73.—The Leaf of Genuine Tea.

and 2 to 3 cm. wide. It is short-stemmed, somewhat thick and fleshy, attenuated at the bottom and usually pointed at the top. At a certain height from the base, from a third to a quarter up, the smooth or wavy border becomes peculiarly, though not deeply, serrated in a regular manner, the serrations, which are hook-shaped, continuing to the tip of the leaf. Mature leaves always show these serrations, but they are somewhat obscure in young leaf buds. The latter, however, are rarely found in this country. The veins extend outward from the central rib nearly parallel to each other, but before reaching the border, each bends upward to form a loop with the oneabove.

Foreign leaves, said to be used as adulterants, are those of the willow, poplar, elder, birch, elm, and rose, but the writer has never found any of these in tea. All of them differ materially from the genuine tea leaf, and if foreign leaves are apparent in a sample under

examination, they should be compared with various leaves collected by the analyst for the purpose.

Stems and Fragments.—These, as well as "tea dust," are apparent by an examination of the leaves, opened out in hot water as explained above. The ash of tea stems and dust is abnormally high. The term "lie tea" is applied to an imitation of tea, consisting of fragments, stems, and tea dust, mixed with foreign leaves, mineral matter, gum, etc.

The ash of such "tea" has been found as high as 50%. Such imitations are now almost unknown. Make-weight substances, such as brickdust, iron salts, metallic iron, sand, etc., have been found in tea. If present, they are to be found in the sediment, obtained on shaking out the tea in water.

Added Astringents.—Catechu is sometimes said to be added to tea to give it increased astringency, especially to such tea as has been adulterated by the addition of exhausted tea. Hagar's method for detecting catechu is as follows:

A hot-water extract of the tea (1 to 100) is boiled with an excess of litharge and filtered. To a part of the filtrate, which should be perfectly clear, nitrate of silver is added. If catechu be present, a yellow flocculent precipitate, rapidly becoming dark-colored, is formed. Pure tea treated in like manner gives a gray precipitate.

Spencer * adds, instead of silver nitrate, a drop of ferric chloride to the clear filtrate. With catechu a green precipitate is formed.

As a matter of fact the worst forms of tea adulteration, such as the actual substitution of foreign leaves, once so commonly practiced, are now extremely rare in this country and have been for some years, by reason of the careful system of government inspection in force at the various ports of entry. The greater portion of the tea on our market to-day is genuine, but fraud is practiced to a considerable extent by the substitution of inferior grades for those of good quality. This form of deception is in many cases beyond the power of the analyst to detect, and properly comes within the realm of the professional tea-taster.

Tea Tablets.—Finely ground tea of varying quality is sometimes pressed into tablets, to be used by travelers and campers for preparing a beverage, by simply dissolving in hot water.

The composition of one of these preparations sold under the name of Samovar Tea Tablets, analyzed by the Mass. State Board of Health, is as follows:

Water	8.7
Theine	2.25
Extract	54.4
Ash	5.4
Soluble ash	2.8
Insoluble ash	2.6

^{*} U. S. Dept. of Agric., Div. of Chem., Bul. 13, p. 885.

Microscopical Structure of Tea.—The powdered tea may be examined directly in water-mount. Schimper recommends treating the powdered tea with chloral hydrate or potash lye, to render it more transparent.

By far the most characteristic element is the peculiarly shaped sclerenchyma, or stone cell, st, Fig. 74, entirely unlike anything to be found in other leaves. These cells are very irregular in form, being sometimes star-shaped, sometimes branched, almost always with deeply wrinkled sides,

Fig. 74.—Powdered Tea under the Microscope. × 160. g, end of leaf nerve; p, chlorophyll parenchyma; st, stone cells; h, hairs. The tissues were warmed in potash to render transparent. (After Moeller.)

and often with sharp points. In most foreign leaves such sclerenchyma cells are lacking, but they are abundant in all genuine tea leaves, excepting rarely in the very young leaves, where they are sometimes not fully developed. They are especially numerous in the main vein and in the stem. They may be seen to best advantage in a section of the stem, or midrib, made parallel to the surface of the leaf. To make such a section, soak the leaf first in water, and afterwards dry in alcohol. The interior of the leaf is composed chiefly of ground tissue, having rounded cells full of chlorophyll grains and the fibro-vascular bundles of the veins.

Other important characteristics are the peculiar hair growth on the under epidermis, B, which is apparent in nearly all teas, also crystal rosettes of calcium oxalate, which are nearly always present, even in fragments of tea leaves, but not in all foreign leaves. The peculiar structure of the lower epidermis, B, with its numerous stomata is also to be noted. See Figs. 189 and 190, Pl. XVIII.

COFFEE.

Nature of Coffee.—Coffee is the seed of the Coffee arabica, a tree which, when under cultivation, is not allowed to exceed twelve feet in height, but when wild sometimes reaches a height of twenty feet. It is indigenous in Southern Abyssinia, and was cultivated in Arabia in the sixteenth century, and in the East Indies in the seventeenth, afterward being introduced into the West Indies and South America. The coffeebeans are usually inclosed in pairs in the berry, being plano-convex with their flat sides together but in "pea berry" coffee only a single, rounded bean is present.

When the ripe fruit is gathered, it is first dried and then freed from the hulls, usually by machinery, or, in the West Indies, the green berries are "pulped" or "hulled" under water by a peculiar macerating machine. The raw beans are roasted, and afterwards ground for preparing the infusion.

Brazil furnishes more than half the world's supply of coffee, and nearly 75% of that consumed in the United States.

Composition of Coffee.—Most of the coffee in the retail market is roasted, being sold either in the whole bean or ground. Comparatively little raw coffee is sold at retail.

The constituents of raw coffee, besides water, are, in the order of their comparative amounts, cellulose or crude fiber, fat, protein, caffetannic acid, sugar, caffein, gum, dextrin, and ash. The effect of roasting coffee, besides driving off most of the water, is to caramelize a large part of the sugar, to make the bean less tough and more brittle, and, most important of all, to develop an empyreumatic, oily substance, known as caffeol (C₈H₁₀O₂), to which the characteristic flavor and aroma of coffee are largely due. Caffeol may be obtained by distilling an infusion of roasted coffee, and extracting the distillate with ether. It is a brown oil, almost insoluble in water.

According to Génin, there are in raw coffee small amounts of two essential oils, one soluble in water, the other insoluble. During the roasting, these are partially transformed into the substance caffeol.

The fat in coffee forms a considerable constituent, amounting in some cases to 15%.

Caffetannic Acid (C₁₅H₁₈O₈) is, when pure, a colorless, crystalline compound. It exists in coffee either as a salt of calcium or magnesium, or, according to Payen, as a caffetannate of potassium and caffeine.

The following summary of analyses of coffee of various kinds made by König show in general its composition:

•	Raw	Coffee.	Roasted Coffee.			
	Minimum.	Maximum.	Minimum.	Maximum		
Water.	8.0	12.0	0.4	4.0		
Caffeine.	0.8	1.8	0.8	1.8		
Fat	11.4	14.2	10.5	16.5		
Reducing sugar	5.8	7.8	0.0	1.1		
Cellulose	16.6	42.3	26.3	51.0		
Total nitrogen	1.1	2.2	1.3	2.7		
Ash	3-5	4.0	4.0	5.0		

The change in composition that takes place in roasting coffee is well shown by the following figures, which give the mean of analyses by König of four samples of coffee before and after roasting:

	Raw Coffee.	Roasted Coffee
Water	11.23	1.15
Caffeine	1.21	1.24
Fat	12.27	14.48
Sugar	8.55	0.66
Cellulose	18.17	10.89
Nitrogenous substances	12.07	13.98
Other non-nitrogenous matter	32.58	45.00
Ash	3.92	4-75

COMPOSITION OF THE ASH OF COFFEE.*

Constituents.	Mocha.	Maracaibo.	Java.	Rio.
Sand	1.44	0.72	0.74	1.34
Silica (SiO ₂)	0.88	0.88	0.91	0.69
Ferric oxide (Fe ₂ O ₃)	0.89	0.89	1.16	1.77
Lime (CaO).	7.18	5.06	4.84	4.94
Magnesia (MgO)	10.68	11.30	11.35	10.60
Potash (K ₂ O)	59.84	61.82	62.08	63.60
Soda (Na ₂ O)	0.48	0.44		0.17
Phosphoric acid (P ₂ O ₅)	12.93	13.20	14.09	11.53
Sulphuric acid (SO ₂)	4.43	5.10	4.10	4.88
Chlorine (Cl)	1.25	0.59	0.73	0.48
j	100.00	100.00	100.00	100.00

The following are analyses of common varieties of roasted coffee, also of coffee substitutes and adulterated coffee made by Lythgoe:†

^{*} U. S. Dept. of Agric., Div. of Chem., Bul. 13, p. 904.

[†] An. Rep. Mass. State Board of Health, 1904, p. 320. U. S. Dept. of Agric., Bur. of Chem., Bul. 90, pp. 43-45.

COMPOSITION OF ROASTED COFFEE.

				اند			cc.	linity N/10 d) of				ou ot	
Variety.		Moisture.	Ash.	Water Soluble Ash.	Sand.	Chlorine.	Ash of 1 Gram of Substance.	I Gram of Ash.	Soluble PrOs.	Insoluble P ₂ O _{3.}	Petroleum Ether Extract.	Index of Refraction Extract at 30°.	Total Nitrogen.
Santos	A B C	1.40 1.87 1.31	4.16 4.31 3.80	3.46 3.62 3.00	 	0.023 .023 .019	2.97 3.36 3.35	75-7		0.346 .295 .295	14.58 13.84 13.86	1-4754 1-4754 1-4750	2.26 2.26 2.39
Rico	A B C	1.29 1.26 1.48	4.05 4.06 4.12	3.30 3.27 3.32	.00	.016 .020 .016	3.53 3.72 3.66	87.2 92.0 88.8	3 .333	-337 -351 -328	13.00 13.34 14.12	1.4752 1.4750 1.4760	2.28 2.26 2.33
Rio {	A B C A	1.76 2.34 2.10 2.05	4.06 3.91 3.74 4.05	3.40 3.24 3.06 3.25	.00.	.020 .021 .023	4.16 3.17 3.22	81.	356	.166 .227 .236 .351	13.38 13.71 13.53 14.84	1.4758 1.4753 1.4756 *1.4737	2.14 2.18 2.26 2.28
Mocha {	B C A	2.95 2.40 3.34	3.85 3.80 4.09	3.07 3.00 3.27	.00	.010	3.94 3.26 3.54 3.88	84.	-333	-354 -364 -545	14.47	*I.4743 *I.4740 I.4752	2.00 2.02 2.48
Java {	B	3·35 3·44	4.38 3.96	3.56	.00	.019	3.54	80.8	3 .194 5 -235	.388 .383	12.28 13.54	1.4758 1.4752	2.35
Highest Lowest		3-44 1.26	4-38 3-74	3.62 3.00	.00	.023	2.95			-545 -166	15.18	1.4760	2.50
Average		2.16	4.03	3.26	.00	.018				-329	13-75	1-4754	2.27
	!		 !			<u> </u>	1	<u> </u>				<u> </u>	
Variety.		r Extract.	tract.	Sugars.		Diastase.			avity -	Re- fer at 20°.	r Cent I	1	
Variety.		Cold Water Extract.	Alcohol Extract.	Reducing Sugars.		Starch by Diastase.	Crode Fiber.	Caffeine.	Specific Gravity at 15°.	Immersion Refractometer Reading at 20°.		1	Ash.
Variety. Santos {	A B C	Cold Water Extract.	16.8	3 0.5	52 2. 58 I.	28 I	Crode Riber.	Caffeine.	1010.1 Specific Gravity at 15°.	O O O C Reading at 20°.		70 2.64 77 2.66 43 2.46	0.40 0.39 30
	B C A B C	20.86 22.72 21.76 22.48 21.76 24.44	16.8 17.1 17.8 15.7 16.3 16.9	3 0.5	52 2. 58 I. 75 2. 50 2. 53 I. 54 2.	28 I 00 I 32 I 17 I 58 I 62 I	3.41 1.02 4.71 3.11 2.93 2.50	1.25 1.10 1.20 1.38 1.21 1.32	1.0107 1.0108 1.0101 1.0107 1.0104	Immersion Re- fractometer 6.95 6.95 Reading at 20.	Index of Re- 1.337 1.337 1.337 1.337 1.338	70 2.64 77 2.66 43 2.46 66 2.60 54 2.50 94 2.77	0.40 -39 -30 -37 -36 -30
Santos {	B C A B C A B C	20.86 22.72 21.76 22.48 21.76 24.44 22.66 22.61	16.8 17.1 17.8 15.7 16.3 16.9 17.0 17.3	3 0.5 1 .0 0 .7 0 .6 6 .0 4 .7	52 2. 58 1. 75 2. 50 2. 53 1. 54 2. 58 2. 78 1. 51 2.	28 II 00 II 32 II 17 II 58 II 62 II 82 II 47 II	3.41 1.02 4.71 3.11 2.93 2.50 4.08 3.10 1.91	1.25 1.10 1.20 1.38 1.21 1.32 1.11 1.10	1.0107 1.0108 1.0101 1.0107 1.0104 1.0103 1.0103	Immersion Re- fractometer 0.95 95 95 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	I . 337 I . 337 I . 337 I . 337 I . 337 I . 337 I . 337	70 2.64 777 2.66 64 2.60 64 2.50 04 2.77 24 2.48 35 2.46 43 2.46	0.40 -39 -30 -37 -36 -30 -40 -36
Santos { Porto { Rico {	B C A B C A B	20.86 22.72 21.76 22.48 21.76 24.44 22.66 22.61 22.73 24.00 20.27	16.8 17.1 17.8 15.7 16.3 16.9 17.0 17.3 17.3 18.0 7 17.9 8 19.5	3 0.13 1 .60 6 .60 6 .60 7 .60 6 .60 6 .60	52 2. 58 1. 75 2. 53 1. 54 2. 58 2. 78 1. 51 2. 78 2. 94 1.	28 I 00 I 32 I 17 I 58 I 62 I 47 I 62 I 30 I 85 I 90 I	3.41 1.02 4.71 3.11 2.93 2.50 4.08 3.10 1.91 1.22 2.34 3.20	1.25 1.10 1.20 1.38 1.21 1.32 1.11 1.10 1.17 1.16 1.10	1.0107 1.0108 1.0101 1.0107 1.0104 1.0113 1.0103	Immersion Refraction et al. 1982	1.337 1.337 1.337 1.337 1.337 1.337 1.337 1.337 1.337	70 2.64 77 2.66 43 2.46 66 2.60 04 2.77 24 2.48 35 2.46 43 2.46 54 2.47 93 2.72	0.40 -39 -30 -37 -36 -30 -40 -36 -36 -40 -36 -36
Santos { Porto Rico } Rio { Mocha { Java	BCABCABCABC	20.86 22.72 21.76 22.48 21.76 24.44 22.66 22.75 24.06 20.27 24.18 23.85 22.16 23.85 22.16	16.8 17.1 17.8 15.7 16.3 16.3 17.0 17.3 17.3 18.0 18.0 17.9	3 0 . 1 . 1 . 1 . 1 . 1 . 1 . 1 . 1 . 1 .	52 2. 58 1. 75 2. 50 2. 53 1. 54 2. 58 2. 78 1. 51 2. 78 2. 54 1. 54 2. 56 3.	28	3.41 1.02 4.71 3.11 2.93 2.50 4.08 3.10 1.91 1.22 2.34 3.20 3.43 3.77 4.75	1.25 1.10 1.20 1.38 1.21 1.32 1.11 1.10 1.17 1.16 1.16 1.18 1.34 1.30	1.0107 1.0108 1.0101 1.0107 1.0103 1.0101 1.0101 1.0101 1.0101 1.0101 1.0101 1.0101 1.0101	Immersion Re- fractometer (190,000) 100,000 10	1.337 1.337 1.337 1.337 1.337 1.337 1.337 1.337 1.337 1.337	70 2.64 77 2.66 43 2.46 65 2.50 04 2.77 24 2.48 35 2.46 43 2.65 58 2.65 58 2.67 22.63 62 2.63	0.40 -39 -30 -37 -36 -30 -40 -36 -40 -39 -38 -38
Santos { Porto Rico } Rio { Mocha { Java	BCABCABCABC.	20.86 22.72 21.76 22.48 21.76 24.44 22.66 22.61 22.70 24.08 24.08 24.08 24.18 23.89 22.19	16.8 17.1 17.8 15.7 16.3 16.3 17.3 17.3 17.3 18.0 17.9 19.5 19.5 19.5 19.5 19.5 19.5 19.5 19	3 0.53	52 2. 58 1. 75 2. 50 2. 53 1. 54 2. 58 2. 78 1. 57 2. 57 2. 58 2. 78 1. 51 2. 52 2. 53 3. 54 3. 57 3. 58	28	3.41 1.02 4.71 3.11 2.93 2.50 4.08 3.10 1.91 1.22 2.34 3.20 3.43	1.25 1.10 1.20 1.38 1.21 1.32 1.11 1.10 1.17 1.16 1.10 1.18 1.34 1.30	1.0107 1.0108 1.0101 1.0107 1.0103 1.0101 1.0101 1.0101 1.0101 1.0101 1.0101 1.0101	Immersion Re- fractometer 26.90.00 26.00.00 27.50.00 26.00.00 27.0	1.337 1.337 1.337 1.337 1.337 1.337 1.337 1.337 1.337 1.337 1.337	70 2.64 77 2.66 43 2.46 66 2.60 04 2.77 24 2.48 35 2.46 43 2.46 54 2.47 77 2.63 66 2.65 66 2.69	0.40 -39 -30 -37 -36 -30 -40 -36 -40 -39 -38
Santos { Porto Rico { Rio { Mocha { Java { Highest	BCABCABCABC	20.86 22.72 21.76 22.48 21.76 24.44 22.66 22.79 24.00 20.27 24.18 23.88 23.88 22.19 23.26	16.88 17.17.88 15.77 16.99 17.00 17.33 17.	3 0.13	52 2. 58 1. 75 2. 50 2. 53 1. 54 2. 58 2. 78 1. 51 2. 78 2. 54 2. 56 3. 78 3. 57 3. 58 3. 59 3. 50 5. 50 5. 50 5. 50 5. 50 5. 50 5. 50 5. 50 5. 50 5. 50 50 5. 50 50 5. 50 50 5. 50 50 5. 50 5	28	3.41 1.02 4.71 3.11 2.93 2.50 4.08 3.10 1.91 1.22 2.34 3.20 3.43 3.77 4.75	1.25 1.10 1.20 1.38 1.21 1.32 1.11 1.10 1.17 1.16 1.16 1.18 1.34 1.30 1.27	1.0107 1.0108 1.0101 1.0103 1.0103 1.0101 1.0101 1.0101 1.0101 1.0101 1.0101 1.0101 1.0101	Immersion Re- fractionneter 26.00.00 26	1.337 1.337 1.337 1.337 1.337 1.337 1.337 1.337 1.337 1.337 1.337 1.337	70 2.64 77 2.66 43 2.46 66 2.50 04 2.77 24 2.48 35 2.46 43 2.46 54 2.72 77 2.63 62 2.58 62 2.58 64 2.72 77 2.63	0.40 -39 -30 -36 -30 -40 -36 -40 -39 -38 -38 -38 -38

^{*} Omitted from average.

15

COMPOSITION OF COFFEE SUBSTITUTES AND OF ADULTERATED COFFEE.

						(cc.	linity N/10 id) of		1		n of		
Variety.	Moisture.	Ash.	Water Soluble Ash.	Sand.	Chlorine.	Ash of 1 Gram of Substance.	r Gram of Ash.	Soluble P2Os.	Insoluble P ₂ O ₅ .	Petroleum Ether Extract.	Index of Refraction of		Total Nitrogen.
Roasted wheat. Roasted chicory Coffee and	5.60 5-55	5.71 4-37	2.8		0.080	0.34		0.649	1.460 -314	2.40 .88			1.84
chicory Coffee,chicory	5.08		3.1			" `	7	1 1	-323	8.32		• • •	1.89
and pea hulls	3.04	4.97	4.0	- 24	*. 28.	2.00	65.6	-472	-740	9.50	1.47	¥5	2.17
	نب			1				1	Ten Per	Cent	Extra	ct.	
Variety.	Cold Water Extract.		Alcohol Extract.	Reducing Sugars.	Starch by Diastase.	Crdue Fiber.	Caffeine	Specific Gravity at 15°.	Immersion Re- fractometer Regulation	Index of Re-	fraction at	Solids.	Ash.
Roasted wheat Roasted chicory	25.8 72.9		.72 -39	4.10 19.34	28.58 2.10		∞.o .∞.	1.0307	45.0	1.3	4463	 7 - 44	 o. 26
Coffee and chicory	31.7		.66			14.31	-95	1.0142	30.5	1.3	3915	3.62	.29
and pea hulls	25.0	× 14	- 25	3.∞	3.78	17.87	1.00			1			

^{*} Admixture of salt.

METHODS OF ANALYSIS.

The sample is prepared for analysis by grinding so as to pass a sieve with holes 0.5 mm. in diameter.

Moisture, Ether Extract, Crude Fiber, Protein, and Ash (including total, water-soluble, water-insoluble, acid-insoluble and alkalinity) are determined as in the case of tea (pp. 368 and 369). Starch, Reducing Matters by Acid Conversion, Sucrose, and Reducing Sugars may be estimated by the methods described under cereal products.

Ten Per Cent Extract. (See page 389.)

Caffetannic Acid.—Krug's Method.*—Two grams of the coffee are digested for thirty-six hours with 10 cc. of water, after which 25 cc. of

^{*}U. S. Dept. of Agric., Div. of Chem., Bul. 13, p. 908.

90% alcohol are added, and the digestion continued for twenty-four hours more. The liquid is then filtered, and the residue washed with 90% alcohol on the filter.

The filtrate, which contains tannin, caffeine, fat, etc., is heated to boiling and a boiling concentrated solution of acetate of lead is added, which precipitates out a caffetannate of lead, Pb₃(C₁₅H₁₅O₈₎₂, containing 49% of lead. When this has become flocculent, it is separated by filtration, and washed on the filter with 90% alcohol, till the washings show



Fig. 75.—Coffee. I. cross-section of berry, natural size. Ph outer pericarp; Mh endocarp; Eh spermoderm; Sa hard endosperm; Sp soft endosperm. II. Longitudinal section of berry, natural size; Dis bordered disc; Se remains of sepals; Em embryo. III. Embryo enlarged; cot cotyledon; rad radicle. (TSCHIRCH AND OESTERLE.)

no lead with ammonium sulphide, and afterwards with ether, till free from fat. It is dried at 100° and weighed.

The weight of caffetannic acid is obtained by multiplying the weight of the precipitate by 652, and dividing by 1263.63.

Woodman and Taylor's Modification.*—To 2 grams of finely ground coffee (passing 0.5 mm. sieve), add 10 cc. of water, and shake for an hour in a mechanical shaking device. Add 25 cc. of 90% alcohol and shake again for half an hour. Filter and wash with 90% alcohol. Bring the united filtrate and washings, about 50 cc., to boiling, and add 6 cc. of saturated lead acetate solution. Separate the precipitated lead caffetannate by means of a centrifuge, decanting the supernatant liquid through a tared filter. Repeat the centrifugal treatment twice with 90% alcohol, decanting each time through the filter. Transfer the precipitate to the filter, and wash free from lead. Wash with ether, dry at 100°, and weigh. The weight of the precipitate multiplied by 0.516 gives the weight of caffetannic acid.

^{*} A. O. A. C. Proc. 1908, U. S. Dept. of Agric., Bur. of Chem., Bul. 122, p. 82.

Caffeine.—Hilger and Fricke Method.*—Boil from 5 to 10 grams of coffee with 100 cc. of water, filter, and treat the residue twice more with boiling water. Add to the united filtrates an excess of lead acetate, filter and wash. Pass hydrogen sulphide into the filtrate to remove the excess of lead, filter, wash and evaporate the filtrate to dryness in a Hoffmeister Schälchen with some sand and a little magnesia. Crush the Schälchen and extract with chloroform until exhausted. Dry the extract at 100° C. and weigh. If the caffeine does not appear to be pure, determine nitrogen in it by the Kjeldahl or Gunning method, multiply the amount of nitrogen by 3.464, thus obtaining the amount of pure caffeine.

ADULTERATION OF COFFEE.

According to the U. S. Standard roasted coffee is coffee which, by the action of heat, has become brown and developed its characteristic aroma, and contains not less than 10% of fat and not less than 3% of ash.

Imitation Coffee.—Formerly, artificial coffee-beans containing no coffee whatever, but cleverly molded to imitate the original, were occasionally to be found, mixed with genuine, whole coffee.†

"Coffee pellets" are occasionally sold in bulk to dealers as an adulterant of whole coffee. These do not closely resemble the real berries in appearance, but are approximately of the same size, and are not apparent to the purchaser when the whole coffee is ground at the time of purchase. A sample of these "pellets" examined recently was found to consist of roasted wheat mash, colored with red ocher.

Coloring Coffee Beans.—The practice of treating raw coffee beans in a manner somewhat analogous to the facing of tea leaves has been sometimes practiced, with a view to giving to cheaper or inferior grades the appearance of high-priced coffee. For this purpose various pigments have been employed, such as yellow ocher, chrome yellow, burnt umber, venetian red, Scheele's green, iron oxide, tumeric, indigo, Prussian blue, etc., the coffee beans being first moistened with water containing a little gum, and shaken with the pigment. As a rule such pigments, especially when inorganic, are best sought for either in the ash, or in the sediment obtained by shaking the coffee beans in cold water, using the

^{*} Arch. Pharm. 1885, p.827, U. S. Dept. of Agric., Bur. of Chem., Bul. 107 (rev.), p. 153. † A sample of such imitation whole coffee in the possession of the writer consists almost entirely of roasted wheat molded into beans with difficulty to be distinguished in appearance from those of genuine coffee, so closely do they resemble the original, even to the cleft in the sides. The chaff in the cleft is, however, lacking.

ordinary qualitative chemical methods. Organic coloring matters can be best extracted with alcohol. Prussian blue and indigo are tested for as in the case of tea leaves (p. 375).

Glazing.—This is a more recent form of treatment of the whole bean, which consists in coating the beans by dipping in egg or sugar, or a mixture of the two, sometimes using various gums. Such glazing is alleged to improve the keeping qualities of the coffee, as well as to aid in clarifying the infusion, and if this is the sole purpose, the practice cannot be condemned as a form of adulteration. If, however, it is done to give inferior varieties of coffee a better appearance, in order to deceive the consumer, it clearly constitutes adulteration within the meaning of the law.

Adulterants of Ground Coffee.—Of the adulterants used in ground coffee the following have been found in Massachusetts: Roasted peas, beans, wheat, rye, oats, chicory, brown bread, pilot bread, charcoal, red slate, bark, and dried pellets, the latter consisting of ground peas, pea hulls, and cereals, held together with molasses.

Methods of Detecting Adulterants.—These methods are, as a rule, physical rather than chemical. A rough test of the genuineness of ground coffee consists in shaking some of the sample in cold water. Pure coffee, under these conditions, usually floats on the surface, while the ordinary adulterants, such as cereals, chicory, mineral ingredients, etc., sink, the grains of chicory coloring the water a brownish-red as they subside.

Macfarlane recommends the use of a saturated solution of common salt, in which a portion of the suspected sample, divided in small grains, is shaken in a test-tube. If the liquid is colored pale amber, while all or nearly all the material floats, the coffee is pure. Any considerable sediment at the bottom of the tube, accompanied by a dark-yellow to brown color imparted to the liquid, indicates adulteration by roasted cereals, or chicory, or both.

A careful examination of the coarsely crushed grains of a ground sample with the naked eye will often serve to detect, and in some cases identify, certain adulterants, such as chicory and ground peas or beans. A magnifying-glass will aid in such an examination, and the observer can often separate the various ingredients of a coffee mixture, first spreading a small portion of the sample on a sheet of white paper. The chicory grains are apparent from their dark and somewhat gummy appearance, and can usually be recognized by crushing them between the teeth. Their soft consistency and sweetish bitter taste are very distinctive. The dull

outer surface of the crushed coffee grains is in marked contrast to the polished appearance of the surface of the broken peas or beans, often to be found as adulterants, while fragments of broken cereal grains are readily distinguished from coffee with a low-power magnifier, though perhaps not easily identified by the eye alone.

Determination of Added Starch.—Starch is determined in the finely powdered sample as directed on page 283.

Microscopical Examination of Coffee.—By far the best means of detecting adulteration is furnished by the microscope. The individual grains of coarsely ground coffee and adulterants, separated by the cold water test or by picking over the mixture, are identified by microscopic examination either after sectioning with a razor or crushing to a powder. In addition, examination is made of a small portion of the sample pulverized in a mortar to a degree fine enough to allow the cover-glass to lie flat on the wetted powder, yet not so fine that it ceases to feel granular when rubbed between the fingers. The writer finds it sufficient to examine this powder in water without further treatment, although Schimper recommends maceration for twenty-four hours with ammonia, in order to render the tissues more transparent, using this reagent also as a mountant.

In general the interior of the coffee tissue or endosperm consists of polygonal cells with highly characteristic, knotty, thickened walls, which are best seen in razor sections, Fig. 76, 2. These cells contain brilliant, colorless, spherical oil drops, and also proteins.

The seed coat is also very characteristic, showing in the powder as occasional delicate silver-like patches, with peculiar, spindle-shaped, thick-sided cells, some of which are loosened from the tissue.

Plates XIV and XV illustrate photomicrographs of pure and adulterated coffee. Fig. 174 shows genuine coffee, with its loose mesh of irregularly polygonal cells, thick-walled, and inclosing oil drops with amorphous material. It is not to be expected that every pulverized sample of genuine coffee, mounted as above, will show in every microscopic field the even, continuous structure that Fig. 174 illustrates, but careful examination will show in nearly every field fragments, and more or less disjointed portions of the polygonal cells, grouped in the form so characteristic of coffee. See Fig. 176.

Chicory under the Microscope.—Fig. 77, after Moeller, shows structural features of chicory. The most striking elements are the fine, thickwalled, long-celled, parenchyma of the bark rp and bp with its delicate

Fig. 75.—Powdered Coffee under the Microscope. ×125. (After Moeller.) 1, seed coat (surface). 2, endosperm parenchyma.

sch 8

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Fig. 77.—Chicory Root in Tangential and Radial Sections. × 160. g, reticulated ducts with perforations qu; hp, wood parenchyma; l, wood fibers; rp, bark parenchyma; sch, milk ducts; bp, bast parenchyma; m, medullary rays. (After Moeiler.)

tracery, and the vessels or ducts g of the wood fibers. These ducts are tubular, resembling jointed cylinders, often with overlapping joints. Less distinct, but very characteristic of certain roots of the composite family, are the narrower branching milk ducts sch which do not exist in beets, turnips, and other roots sometimes substituted for chicory.

Fig. 178, Pl. XV, is a photomicrograph of an adulterated sample of coffee, showing in this particular field chicory alone. It is a mass of confused cellular tissue, traversed by two broad bands of the vessels, with their striking, transverse, dotted markings.

Fig. 177, Pl. XV, shows a sample of coffee adulterated with roasted peas and pea hulls. No genuine coffee appears in this field. The chief masses in the center are characteristic aggregations of the round starch granules of the roasted pea. The rectangular billets, like bunches of matches, are from the outer or palisade layer of the pea.

Fig. 164, Pl. XI, and Fig. 154, Pl. IX, show the close resemblance between the starches of the pea and bean, both of which are commonly used in coffee.

The palisade structures of the hulls of these legumes also bear a close resemblance, but the cells of the next layer in the pea are hour-glass shaped, while in the bean they are not remarkable for their shape, but for the single crystal of calcium oxalate contained in each.

The effect of roasting on starches used as adulterants of coffee is to twist and distort the granules, in some cases destroying largely the even structure of the raw starch. Starch granules of wheat, barley, and rye, for example, are almost perfect circular disks in the case of the raw starch, while in roasted products, such as pilot biscuit and stale bread, the granules are twisted and distorted, sometimes almost forming the letter "S."

Use of Chicory in Coffee.— Chicory is a perennial herb (Cicorium intybus) of the same family (Compositæ) as the dandelion. The roasted and pulverized chicory root is so much used in ground coffee to impart a peculiar flavor thereto, that by many it is considered as not strictly an adulterant. The taste imparted to coffee by a small admixture of pure chicory is to some desirable, but if its unrestricted use is sanctioned in this manner, the door would soon be opened to a more unlimited form of adulteration, wherein the chicory might predominate. It is, therefore, best to regard chicory as an adulterant, and to require the package containing a mixture of coffee and chicory, if sold legally, to have plainly printed thereon the percentage of chicory in the mixture.

Chicory, when roasted, consists of gum, partly caramelized sugar, and insoluble vegetable tissue. Common adulterants of chicory are dried beets and other roots, also cereal matter.

Villiers and Collin * give the following analyses of two samples of chicory:

		In Large Granules.	In Powder.
Soluble in water:	Water (loss at 100° to 103°). Weight of total matter soluble in water. Reducing sugar Dextrin, gum, inulin. Albuminoids. Mineral matter. Coloring matter. Albuminoids.	57.96 26.12 9.63 3.23 2.58 16.40	16.96 56.90 23.79 9.31 3.66 2.55 17.59 2.98
Insoluble in water:	Weight of the total insoluble matter. Mineral matter. Fat Cellulose.	25.76 4.58	26.14 5.87 3.92 13.37

See also analysis of roasted chicory on page 382.

Detection and Estimation of Chicory.—Various chemical tests for detection of chicory in coffee infusions have been suggested, depending on color reactions,† but these are, as a rule, unreliable. By far the best means for detecting chicory in coffee is furnished by the microscope.

In mixtures containing coffee and chicory only, the approximate amount of the latter can be calculated from the specific gravity of a 10% decoction, using conveniently the method of McGill.‡ A quantity of the pulverized sample, corresponding to 10 grams of the dry substance, is weighed in a counterbalanced flask, and water added till the weight of the contents is 110 grams. Fit the flask with a reflux condenser, and after so regulating the heat that boiling begins in ten to fifteen minutes, continue the boiling for an hour. Remove the flame, and after fifteen minutes pass through a dry filter, cool, and determine the specific gravity at 15°. McGill found the average specific gravity of a 10% decoction as above carried out to be, in the case of pure coffee, 1.00986 and in the case of chicory 1.02821, the difference being 0.01835.

The specific gravity of the 10% decoction of the suspected sample

^{*} Falsifications et Alterations des Substances Alimentaires, p. 234.

[†] See Allen's Commercial Org. Analysis, Vol. III, pt. II, p. 540.

[†] Trans. Royal Soc. of Canada, 1887.

at 15° being d, the per cent of chicory, c, can be calculated roughly by the formula

$$c = 100 - \frac{(1.02821 - d)100}{0.01835}$$

This method is of course inapplicable when other substances than chicory are present.

Date Stones, roasted and ground, have been used to some extent as a coffee adulterant. Fig. 78 shows the structural features of date stones

Fm. 78.—Powdered Date Stones under the Microscope. end, endocarp; e, episperm; e, albumen in cross-section; e', albumen in longitudinal section. (After Villiers and Collin.)

under the microscope. *End* represents a fragment of endocarp with its elongated, thick-walled cells, peculiarly arranged as shown, adjacent cells often lying with axes at right angles to each other. The more evenly formed episperm cells, e, are thin-walled and of a brown color. The albumen, a, is made up of very thick-walled, somewhat regularly arranged cells, indented from within with deep channels. Date stones are readily distinguished from coffee by these features.

Hygienic Coffee.—Various processes have been devised for removing the caffeine from coffee. One of these, patented in Germany, has recently come into extensive use, as the flavor of the beverage is not greatly injured by the treatment. In following out this process the whole beans are first exhausted with water in a vacuum, and the infusion extracted with a suitable solvent for caffeine. The exhausted beans are then impregnated with the decaffeinated infusion and dried in a vacuum. This treatment,

as shown by the investigations of Lendrich and Murdfield,* does not completely remove all the caffeine, the quantity remaining being from 0.14 to 0.26%, or about one-sixth of that in the untreated coffee. Further effects of the treatment are a decrease in the water extract and an increase in the fat. The following are the average of analyses, made by these authors, of caffeine-free and untreated coffee:

			Analysis of the Dry Substance.							
	Number of Analyses.	Moisture.	Ash.	Water Soluble Ash,	Alkalinity of Ash (cc. N/1 HCl per 100 grams of Coffee).	Water Extract.	Fat (Petroleum Ether Extract).	Caffeine (Juckenack and Hilger	Protein (Protein Nitrogen × 6.25).	
"Caffeine-free Coffee" Untreated coffee	14 9	% 2.13 1.46		% 3.22 3.77	47.72 56.43	% 21.30 26.17	% 17.13 15.73	% 0.22 1.19	% 11.83 11.75	

Several brands of coffee, advertised to be free from tannin and in some cases also from caffeine, have been placed on the market in the United States. Some of these consist merely of ground coffee from which the chaff (which is represented to contain not only the tannin but also most of the caffeine) has been removed by mechancial means. The absurdity of the claims of the manufacturers is shown by the following analyses made in New Hampshire by C. D. Howard.†

	Water.	Ash.	Fat.	Fiber.	Caffeine.	Caffe- tannic Acid.
Tanninless coffee No. 1. Tanninless coffee No. 2. Tanninless coffee No. 3. Java and Mocha. Coffee chaff.	2.70	4.10	13.16	18.46	1.17	10.76
	2.70	4.05	14.12	15.70	1.3	11.04
	2.26	3.61	12.55	22.70	0.87	7.61
	3.13	4.13	14.10	15.50	1.29	11.17
	2.60	5.65	9.30	26.50	0.40	5.98

The following analyses made at the Connecticut Station by E. J. Shanley‡ corroborate those of Howard:

^{*} Zeits. Unters. Nahr. Genuss., 15, 1908, p. 705.

[†] A. O. A. C. Proc. 1906, U. S. Dept. of Agric., Bur. of Chem., Bul. 105, p. 41.

[‡] An. Rep. Conn. Exp. Sta., 1907, p. 141.

	Caffeine in the Coffee.	Caffetannic Acid in the Coffee.	Caffetannic Acid in the Chaff.	Per Cent of Chaff in the Coffee.
Tanninless coffee A	1.11	9.89 9.45 9.96		
Tanninless coffee C	1.13	9.90 9.51 9.96	5.46 7-55	1.80 2.38
Rio coffee	1.13	9-47	7-55 6.79	1.77

Coffee Substitutes.—A large number of preparations sold as "coffee substitutes" or "cereal coffee" are now on the market in the United States, most of which are composed, as alleged on the labels, of cereals, ground peas, etc. Some contain roasted wheat, malt or some other cereal alone, others are mixtures of cereals or cereal products and peas, and a few contain chicory. Some of these preparations have labels calling attention to the evil effects of coffee, and one of the latter class, extensively advertised, and purporting to contain nothing but the entire wheat kernel roasted and ground, was found to contain peas, and about 30% of that "most harmful ingredient" coffee itself. Various substitutes are also made from dried fruits such as figs, prunes and bananas.

In addition to the materials named the following have been used in Europe: beans, lupine seeds, cassia seeds, astragalus seeds, Parkia seeds, chick peas, soja beans, dried pears, carob bean pods, date stones, ivory nuts, acorns, grape seeds, fruit of the wax palm, cola nuts, false flaxseed, dandelion roots, beets, turnips and carrots.*

As in the case of coffee the analyst must depend chiefly on the microscope in identifying the constituents of coffee substitutes. Coffee itself should properly be considered in the light of an adulterant.

COCOA AND COCOA PRODUCTS.

Nature of the Cocoa Bean.—The various chocolate and cocoa preparations are made from the bean of the tree *Theobroma cacao*, of the family of *Byttneriacea*. This tree averages 13 feet in height, and its main trunk is from 5 to 8 inches in diameter. It is a native of the American tropics, being especially abundant and growing under best conditions in Mexico, Central America, Brazil, and the West Indies.

The cocoa beans of commerce are derived chiefly from Ariba, Bahia, Caracas, Cayenne, Ceylon, Guatemala, Haiti, Java, Machala, Mara-

^{*} Winton's Microscopy of Foods, p. 435.

caibo, St. Domingo, Surinam, and Trinidad. Besides these, the Seychelles and Martinique furnish a small amount.

The plant seeds, or beans, grow in pods, varying in length from 23 to 30 cm., and are from 10 to 15 cm. in diameter. The beans, which are about the size of almonds, are closely packed together in the pod. Their color when fresh is white, but they turn brown on drying.

The gathered pods are first cut open, and the seeds removed to undergo the process of "sweating" or fermenting, which is carried out either in boxes or in holes made in the ground. This process requires great care and attention, as upon it depends largely the flavor of the seed. The sweating operation usually takes two days, after which the seeds are dried in the sun till they assume their characteristic warm red color, and in this form are shipped into our markets.

Manufacture of Chocolate and Cocoa.—For the production of chocolate and cocoa the beans are cleaned and carefully roasted, during which process the flavor is more carefully developed, and the thin, paper-like shell which surrounds the seed is loosened, and is very readily removed. The roasted seeds are crushed, and the shells, which are separated by winnowing, form a low-priced product, from which an infusion may be made, having a taste and flavor much resembling chocolate.

The crushed fragments of the kernel or seed proper are called *cocoa nibs*, and for the preparation of chocolate they are finely ground into a paste and run into molds, either directly, or after being mixed with sugar and vanilla extract or spices, according to whether plain or sweet chocolate is the end product.

For making cocoa, however, a portion of the oil or fat known as the cocoa butter is first removed, by subjecting the ground seed fragments to hydraulic pressure, usually between heated plates, after which the pressed mass is reduced to a very fine powder, either directly, or by treatment with ammonia or alkalies, to render the product more soluble. It is held that the large amount of fat contained in the cocoa seeds (varying from 40 to 54 per cent) is difficult of digestion to many, such as invalids and children, and hence the desirability of removing part of the fat.

Composition of Cocoa Products.—The chief constituents of the raw cocoa bean, named in the order of their relative amount, are fat, protein, starch, water, crude fiber, ash, theobromine, gum, and tannin. During the roasting there is reason to believe a volatile substance is developed much in the nature of an essential oil, which gives to the

product its peculiar flavor, and is somewhat analogous to the caffeol of coffee.

Tannin, the astringent principle of cocoa, exists as such in the raw bean, but rapidly becomes oxidized to form cocoa red, to which the color of cocoa is due.

Weigmann gives the following results of analyses of cocoa nibs and shells:

COMPOSITION OF COCOA NIBS.

Commercial Varieties.	Moisture.	Nitrogenous Substances, including Theobromine.	Theobromine.	Fat.	Starch.	Other Car- bohydrates.	Cellulose.	Ash.	Sand.
Caracas. Trinidad. Surinam Port au Prince. Machata Puerto Cabello. Ariba.	7.77 7.87 7.53 7.77 8.17 8.08 8.27	14.56 14.06 13.50	1.31	45.54 44.62 44.74 46.35 45.93 46.61 45.15	25.30 26.45 5.97 5.69	15.53 17.50 16.96	4.36	4.91 3.48 3.16 4.15 4.09 4.28 3.88	2.06 0.10 0.13 1.48 0.22 0.18

COMPOSITION OF COCOA SHELLS.

Commercial Varieties.	Moisture.	Nitrogenous Substances.	Theobromine.	Fat.	Nitrogen-free Extract.	Cellulose.	Ash.	Sand.	Total Nitro-gen.
Caracas. Trinidad. Surinam Puerto Cabello.	12.49 14.64 13.93 14.89	16.25	0.74	2.38 3.45 2.54 2.01	44.89	15.79	6.19 6.63	6.26 0.42 0.85 0.27	2.11 2.34 2.60 2.59

The following are the summarized results of the analyses of seventeen varieties of cocoa seeds and shells, made by Winton, Silverman, and Bailey.*

^{*} An. Rep. Conn. Agric. Exp. Sta., 1902, p. 270.

	Roasted Cocoa Nibs.							
	Air	-dry Mate	erial.	Wate	Water- and Fat-free Material.			
	Maxi- mum.	Mini- mum.	Mean.	Maxi- mum.	Mini- mum.	Mean.		
Water	3.18	2.29	2.72					
Total ash		2.61	3.32	18.8	5.76	7.04		
Water-soluble ash	1.86	0.73	1.16	3.96	1.60	2.46		
Ash insoluble in acid	0.07	0.00	0.02	0.14	0.00	0.05		
Alkalinity of ash	3-35	1.50	2.51	7.12	3.29	5.32		
Theobromine	1.32	0.82	1.04	2.92	1.66	2.27		
Caffeine	0.73	0.14	0.40	1.55	0.31	0.86		
Other nitrogenous substances	13.06	11.00	12.12	28.05	23.37	25.69		
Crude fiber		2.21	2.64	6.56	4.70	5.61		
Crude starch (acid conversion)	12.37	9.30	11.16	25.68	19.80	23.66		
Pure starch (diastase conversion)	8.99	6.49	8.07	18.61	13.82	17.10		
Other nitrogen-free substances		17.69	19.57	44.08	38.78	41.49		
Fát	52.25	48.11	50.12		• •			
Total nitrogen		2.20	2.38	5.41	4.74	5.05		
Constants of fat (ether extract):		1		_		" "		
Melting-point, degrees C	35.0	32.3	33-3					
Zeiss refractometer reading at 40° C	48.00	46.00	47.23					
Refractive index at 40° C	1.4579	1.4565	1.4573					
Iodine number	37.89	33.74	34-97					
Per cent of nibs in whole bean	92.90	86.12	88.46					
" " shells " "	13.88	8.83	11.54					

	Roasted Cocoa Shells.							
	Air-	dry Mate	rial.	Wate	ater- and Fat-free Material.			
·	Maxi- mum.	Mini- mum.	Mean.	Maxi- mum.	Mini- mum.	Mean.		
Water	6.57	3.71	4.87	'				
Total ash	20.72	7.14	10.48	21.97	5.63	11.33		
Water-soluble ash	5.67	2.02	3.67	6.11	2.16	3.97		
Ash insoluble in acid	11.18	0.05	2.51	11.86	0.05	2.70		
Alkalinity of ash	5.92	5.02	5.52	6.47	5-32	5-97		
Theobromine	0.90	0.20	0.49	0.97	0.22	0.52		
Caffeine	0.28	0.04	0.16	0.31	0.04	0.17		
Other nitrogenous substances	18.06	10.69	14-54	19.40	11.34	15.70		
Crude fiber	19.21	12.93	16.63	20.72	13.71	18.01		
Crude starch (acid conversion)	13.89	9.87	11.62	15.42	10.47	12.59		
Pure starch (diastase conversion)	5.16	3.36	4.14	5-59	3.65	4.47		
Other nitrogen-free substances	51.86	43.71	46.40	55.84	47.04	50.08		
Fat	5.23	1.66	2.77			1		
Total nitrogen	3-17	1.74	2.34	3-41	1.87	2-54		

According to Bell * the ash of cocoa nibs has the following composition:

	Per Cent.
Sodium chloride	0.57
Soda	0.57
Potash	. 27.64
Magnesia	
Lime.	
Alumina	. 0.08
Ferric oxide	. 0.15
Carbonic acid	. 2.92
Sulphuric acid	• 4-53
Phosphoric acid	
	T00.00

100.00

Theobromine (C₇H₈N₄O₂), the chief alkaloid of cocoa, when pure, forms a white, crystalline powder, having a bitter taste. It is slightly soluble in water and alcohol, very slightly soluble in ether, insoluble in petroleum ether, but readily soluble in chloroform. It sublimes at 290° to 295° C. It is a weak base, and much resembles caffeine. A small amount of caffeine has also been found in cocoa, but in most analyses is reckoned in with the theobromine.

The Nitrogenous Substances of Cocoa, aside from the alkaloids, have been little studied. Stutzer has, however, separated them roughly as in the following analyses of four samples, of which A was manufactured without chemicals, B with potash, and C and D with ammonia:

	A.	В.	c.	D.
Total nitrogen	3.68	3.30	3.95	3-57
Theobromine	1.92	1.73	1.98	1.80
Ammonia	0.06	0.03	0.46	0.33
Amido compounds	1.43	1.25	0.31	
Digestible albumin	10.25	7.68	10.50	1.31 7.81
Indigestible nitrogenous substances	7.18	9.19	7.68	8.00
Containing nitrogen	1.15	1.47	1.23	1.28
Proportion of total nitrogen indigestible.	31.2	44-5	31.2	35.8

Pentosans.—Several authors have called attention to the value of these substances as a means of detecting added shells in cocoa products. Lührig and Segin† found in cocoa nibs from 2.51 to 4.58 per cent

^{*} Analysis and Adulteration of Foods.

[†] Zeits. Unters. Nahr. Genuss., 12, 1906, p. 161.

of pentosans calculated to the dry, fat-free substance, and in the shells from 7.59 to 11.23 per cent calculated to the dry substance.

Milk Chocolate, a product of comparatively recent introduction, consists of a mixture of chocolate, sugar, milk powder, and cocoa butter. It is especially prized by travelers and others who desire a concentrated, and at the same time palatable food.

The following analyses by Dubois* show the composition of three of the leading brands on the market, and also illustrate the accuracy of Dubois' method of determining sucrose and lactose given on page 399.

	Polarization.				Su-	Lac-		Approx. Per Cent
	Direct.	After Inver- sion.	Temp. °C.	At 86°.	Crose, Per Cent.	tose, Per Cent.	Meissl Num- ber of Fat.	Butter Fat in Total Fat.
Commercial milk chocolate:	,			1				
A	+21.00	- 2.00	24	+1.36	40.90	8.24	5-3	22.1
В	+ 23.22	-2.22	23	+1.50		9.12	5.5	22.Q
C	+ 23.88			+1.36			5.8	24.2
Milk chocolate made in the						=	*	!
laboratory:							i	
D { Found	+19.00	-1.50	20	+1.40	35-99	8.52	4.83	20.1
Calculated					35.82			
E Found	+19.70	-2.20	21	+0.99		6.03	3.48	14.5
Calculated					39.80	5.88		
,				!		_		

Various Compounds of chocolate or cocoa with other materials have been placed on the market. Zipperer † gives formulas or analyses of seventy-four such preparations, containing one or more of the following ingredients: oatmeal, barley meal, malt, malt extract, wheat flour, potato flour, rice, peas, peanuts, acorns, cola nuts, sago, arrowroot, Iceland moss, gum Arabic, salep, dried meat, meat extract, peptones, milk powder, plasmon (a preparation of casein), eggs, saccharin, vanilla, spices, and inorganic salts. Certain medicinal preparations also contain cocoa products.

Cocoa Butter.—See page 529.

^{*} Jour. Am. Chem. Soc., 29, 1907, p. 556.

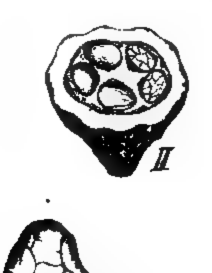
[†] The Manufacture of Chocolate and Cacao Preparations, 2d ed., 1902.

METHODS OF ANALYSES.

Preparation of the Sample.—Cocoa is usually in a fine powder, and needs merely to be put through a sieve, to break up lumps, and mixed. Chocolate should be grated or shaved so as to permit mixing. It can not be ground, as the heat of grinding reduces it to a paste.

Moisture.—Dry two grams of the material to constant weight at 100° C, in a current of dry hydrogen. Somewhat lower results are obtained by drying in a dish in air.

Ash.—Proceed as described under tea (page 369) in the determination of total, water-soluble and acid-soluble ash, and the alkalinity of the ash.



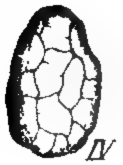




Fig. 79.—Cocoa. I entire fruit, $\times \frac{1}{4}$; II fruit in cross-section; III seed (cocoa bean) natural size: IV seed deprived of seed coat; V seed in longitudinal section, showing radicle (germ); VI seed in cross-section. (Winton.)

Protein.—Determine total nitrogen by the Kjeldahl or Gunning method. From the percentage of total nitrogen subtract the nitrogen of the theobromin and caffeine, obtained by multiplying the percentages found by 0.311 and 0.289 respectively, and multiply the remainder by 6.25.

Fat (Ether Extract).—Extract two grams of the material in a continuous extractor until no more fat is removed. Grind the residue and repeat the extraction. Dry the combined extract at 100° C. and weigh.

Constants of Fat. - See chapter on Edible Oils and Fats.

Crude Fiber.—Proceed as in the analysis of cereal products (page 277), using the residue from the ether extraction.

Reducing Matters by Acid Conversion (Crude Starch).*—Weigh four grams of the material into a small Wedgewood mortar, add 25 cc. of ether, and grind with a pestle. After the coarser material has settled out, decant off the ether with the fine suspended matter on a 11 cm. paper. Repeat this treatment until no more coarse material remains. After the ether has evaporated, transfer the fat-free residue from the filter to the mortar by means of a jet of cold water, and rub to an even paste. Filter the liquid on the paper previously employed. Repeat the process of transferring from the filter to the mortar, grinding, and filtering, until all sugar is removed. In the case of sweetened cocoa products, at least 500 cc. of water should be used.

Transfer the residue to a 500-cc. flask by means of 200 cc. of water, and convert the starch into dextrose by Sachsse's method (page 283).

Cool the acid solution, nearly neutralize with sodium hydroxide solution, add 5 cc. of lead sub-acetate solution (page 586), make up to 250 cc. and filter through a dry filter. To 100 cc. of the filtrate, add 1 cc. of 60% sulphuric acid, shake thoroughly, allow to settle, and filter through a dry filter.

Determine reducing matters by Allihn's method (page 608).

Dubois,† instead of treating with ether as above described, shakes four grams of the unsweetened product or eight grams of the sweetened with 100 cc. of gasoline, and whirls in a centrifuge to separate from the insoluble matter. After decanting off the gasoline layer, sweetened products are treated in like manner with two portions of 100 cc. of water to remove the bulk of the sugar, and finally washed on the paper.

Starch.—Diastase Method.—Remove the fat and sugar from four grams of the material by treatment with ether and water, as described in the preceding section, and determine starch in the residue by the diastase method (page 283).

Pentosans. See page 285.

Sucrose and Lactose.—Dubois's Method.‡—Extract the fat from 13 grams of the sample by shaking and centrifuging twice with 100 cc. of gasoline, separating the solvent by decantation. To the residue add 100 cc. of water, and shake for ten minutes. Add 5 cc. of basic lead acetate solution (p. 586), filter, and remove the excess of lead. Allow 25 cc. of this solution to stand over night to destroy birotation, and polarize. Mul-

^{*} Winton, Silverman and Bailey, An. Rep. Conn. Exp. Sta., 1902, p. 275.

[†] A. O. A. C. Proc. 1908, U. S. Dept. of Agric., Bur. of Chem., Bul. 122, p. 214.

[‡] Jour. Am. Chem. Soc., 29, 1907, p. 556.

tiply readings by 2. Invert 50 cc. of the above filtrate as described on page 588, nearly neutralizing the acid after cooling with sodium hydroxide solution, and make up to 100 cc. Bring to temperature at which direct readings were made and polarize, and also polarize at 86° in a water jacketed tube; multiply all invert readings by 4.

Calculate the approximate weight of sucrose and lactose present in the 13 grams by the following formulas:

Grams of sucrose =
$$\frac{(a-b)1.05}{t} \times 13$$
.

Grams of lactose =
$$\frac{c \times 1.264 \times 1.11 \times 1.05 \times 13}{100} = \frac{19.152c}{100}$$
.

a=direct readings, normal weight.

b = invert readings, normal weight.

c =invert readings, normal weight, at 86° C.

From the total amount of sugar found by the above, obtain the value of x from the table below, and calculate sugars as follows:

$$\frac{(a-b)1.05x}{142.66 - \frac{t}{2}} = \text{per cent of sucrose.} \quad (1.473c)x = \text{per cent of lactose.}$$

2 grams of sugar in sample, x = 101.2

4 grams of sugar in sample, x = 102.5

6 grams of sugar in sample, x = 103.6

8 grams of sugar in sample, x = 104.8

10 grams of sugar in sample, x = 106.05

15 grams of sugar in sample, x = 109.40

20 grams of sugar in sample, x = 112.40

Theobromine and Caffeine (Decker-Kunze Method).*—Boil 10 grams of the powdered material and 5 grams of calcined magnesia for 30 minutes with 300 cc. of water. Filter by the aid of suction on a Buchner funnel, using a round disk of filter paper. Transfer the material and paper to the same flask used for the first boiling, add 150 cc. of water,

^{*} Schweiz. Wchschr. Phar., 40, 1902, pp. 527, 541, 553; Abstract Chem. Centr., 74, 1903, p. 62; An. Rep. Conn. Exp. Sta., 1902, p. 274.

and boil 15 minutes. Filter as before, and repeat the operation of boiling with 150 cc. of water and filtering. Wash once or twice with hot water. Evaporate the united filtrates (with quartz sand if sugar be present), to complete dryness in a thin glass dish of about 300 cc. capacity.*

Grind to a coarse powder in a mortar provided with a suitable cover to prevent loss by flying. Transfer to the inner tube of a continuous fat extractor, and dry thoroughly in a water oven. Extract with chloroform for 8 hours, or until the theobromine and caffeine are completely removed, into a weighed flask. It is important that the material be thoroughly dry, that an extractor be used that permits of a hot extraction, and that a considerable volume of chloroform passes through the material. Distil off the chloroform, and dry at 100° C. to constant weight.

If the material be pure chocolate or cocoa, the extract thus obtained is practically pure theobromine and caffeine, but if the material is cocoa shells or a cocoa product mixed with a large amount of shells, the extract may be brown in color, due to the presence of considerable amounts of impurities.

In either case, separate the caffeine by treating the extract in the flask at the room temperature for some hours with 50 cc. of pure benzol. Filter through a small paper into a tared dish, evaporate to dryness, and dry to constant weight at 100° C., thus obtaining the amount of caffeine.

Determine theobromine by Kunze's † method, as follows:

Add to the residue and paper 150 cc. of water, enough ammonia water to make the liquid slightly alkaline, and an excess of decinormal silver nitrate solution. Boil to half the original volume, add 75 cc. of water, and repeat the boiling. The solution should be perfectly neutral. If it contains the slightest amount of free ammonia, add water and boil until it is completely removed.

Filter from the insoluble silver theobromine compound, and wash with hot water. In the filtrate determine the excess of silver nitrate by Volhard's ‡ method as follows:

Add 5 cc. of cold saturated solution of ferric ammonium sulphate (ferric-ammonium alum), and enough boiled nitric acid to bleach the liquid. Titrate with decinormal ammonium sulphocyanide solution until a permanent red color appears.

^{*}A "Hoffmeister Schälchen" may be used, or dishes may be made from broken flasks by making a scratch with a diamond and leading a crack from this scratch about the flask by means of a glowing springcoal.

[†] Ztschr f. anal. Chem., 33, 1894, p. 1.

[‡] Ibid., 13, 1874, p. 171.

One cc. of decinormal AgNO₃ solution is equivalent to 0.01802 gram of theobromine. If the mixed alkaloids were colorless, the theobromine obtained by subtracting the weight of caffeine from the weight of the mixed alkaloids will usually agree closely with that obtained by silver titration.

ADULTERATION OF COCOA PRODUCTS AND STANDARDS OF PURITY.

The following are the U. S. standards:* Standard chocolate should contain not more than 3% of ash insoluble in water, 3.5% of crude fiber, and 9% of starch, nor less than 45% of cocoa fat.

Standard sweet chocolate and standard chocolate coating are plain chocolate mixed with sugar (sucrose), with or without the addition of cocoa butter, spices, or other flavoring material, containing in the sugarand fat-free residue no higher percentage of either ash, fiber, or starch than is found in the sugar- and fat-free residue of plain chocolate.

Standard cocoa should contain percentages of ash, crude fiber, and starch corresponding to those of plain chocolate, after correcting for fat removed.

Standard sweet cocoa is cocoa mixed with sugar (sucrose) containing not more than 60% of sugar, and in the sugar- and fat-free residue no higher percentage of either ash, crude fiber, or starch than is found in the sugar- and fat-free residue of plain chocolate.

The removal of fat, or the addition of sugar beyond the above prescribed limits, or the addition of foreign fats, foreign starches, or other foreign substances, constitutes adulteration, unless plainly stated on the label.

The most common adulterants of cocoa are sugar and various starches, especially those of wheat, corn, and arrowroot. Starch is sometimes added for the alleged purpose of diluting the cocoa fat, instead of removing the latter by pressure, thus, it is claimed, rendering the cocoa more digestible and more nutritious. Unless its presence is announced on the label of the package, starch should be considered as an adulterant. Cocoa shells are also commonly employed as a substitute for, or an adulterant of, cocoa. Other foreign substances found in cocoa are sand and ground wood fiber of various kinds. Iron oxide is occasionally used as a coloring matter, especially in cheap varieties.

^{*} U. S. Dept. of Agric., Off. of Sec., Circ. 19.

Such adulterants as the starches and cocoa shells are best detected by the microscope. The presence of any considerable admixture of sugar is made apparent by the taste. Mineral adulterants are sought for in the ash.

Addition of Alkali.—The amount of water-soluble matter in cocoa is very small (about 20 to 25 per cent), and in preparing the beverage, the desideratum aimed at is to produce as perfect an emulsion as possible. The legitimate means of accomplishing this is by pulverizing the cocoa very fine, so that particles remain in even suspension and form a smooth paste. Another means sometimes resorted to for producing a so-called "soluble cocoa" is to add alkali in its manufacture, the effect being to act upon a part of the fat, and produce a more perfect emulsion with less separation of oil particles. Such treatment with alkali is regarded with disfavor, even if not considered as a form of adulteration. Cocoa thus treated is generally darker in color than the pure article.

The use of alkali is usually rendered apparent by the abnormally high ash, and by the increased alkalinity of the ash, the latter constant being expressed in terms of the number of cubic centimeters of decinormal acid necessary to neutralize the ash of r gram of the sample. In pure, untreated cocoa, the ash rarely exceeds 5.5%, and the alkalinity of the ash is generally not more than 3.75. In cocoa treated with alkali, the ash sometimes reaches 8.5%, with the alkalinity running as high as 6 or even 8.

Microscopical Structure of Cocoa.—Fig. 80 shows elements of the powdered cocoa bean, both of the shell and of the kernel. The powder of the latter should constitute pure cocoa, with occasional fragments only of the shell. The irregular lobes constituting the kernel are each inclosed in a membrane made up of angular cells, filled with granular matter. (4), (5), and (6) show elements of the powdered cotyledons, or seed kernels. The polygonal tissue of the cotyledon is shown in cross-section at (4). In the powder one finds also dark granular matter, bits of débris, and fragments, with masses of yellow, reddish-brown, and sometimes violet coloring matter, together with numerous starch granules and aleurone grains.

The starch granules are nearly circular, with rather indistinct central nuclei, and range in size from 0.0024 to 0.0127 mm., averaging about 0.007 mm. They are more often found in single detached grains, but sometimes in groups of two or three. Occasional spiral ducts, sp, are seen, but these are not abundant in the pure cocoa.

The masses of color pigment are shown up with striking clearness, according to Schimper, by applying a drop of sulphuric acid to the edge of the cover-glass and allowing it to penetrate the tissue. The bits of coloring matter are for a short time colored a brilliant red, which, however, soon fades. Ferric chloride colors them indigo blue.

Schimper recommends mounting the powder in a drop of chloral hydrate, which soon renders most of the tissues transparent. It is sometimes necessary to allow the chloral to act on the powder in a closed

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Fig. 8o.—Cocoa under the Microscope.

- A. Powdered Cocoa under the Microscope. ×125. (After Moeller) 1, cross-section through shell parenchyma; 2, thick-walled cells; 3, epidermis of shell (surface section); 4, cross-section of cotyledon tissue, 5, 6, cotyledon parenchyma; 7, starch.
- B. Cocoa Shell in Surface Section. \times 160. ϵp , epicarp; p, parenchyma of the fruit; q u, layer of transverse cells. (After Moeller.)

vessel for twenty-four hours, before all the elements, of pure cocoa are rendered transparent. If after that time opaque masses are still found, these are due to foreign material.

Ammonia may be used instead of chloral with even better results, but this reagent requires longer treatment, soaking for several days or a week being sometimes necessary.

Fig. 185, Pl. XVII, shows the microscopical appearance of genuine powdered cocoa with its variously sized starch grains and the débris of the ground cotyledons. Fig. 186 shows cocoa adulterated with arrowroot.

Cocoa Shells.—The shell parenchyma with side view of the stone-cell layer a and frequent spiral ducts, all characteristic of the ground shell, are shown at 1, Fig. 80.

In plan view the thick-walled stone-cell layer is shown at 2, and the spongy, outer seed-skin tissue, composed of two layers, with elongated cells running crosswise to each other in striated fashion, and with the underlying hairs or so-called "Mitscherlich bodies," is shown at 3. The presence of an abnormally large number of yellow and brown fragments in the water-mounted cocoa specimen, even under small magnification, arouses suspicion of the presence of shells, the most distinctive elements of which are the spongy tissue, the stone cells, and the abundant spiral ducts, the latter being scarce in pure cocoa powder.

Cocoa shells are indicated on chemical analysis by the abnormally high ash, crude fiber and pentosans.

Added Starch.—This can only be approximately determined by a careful examination with the microscope. Long experience will enable the analyst to familiarize himself with the appearance and abundance of starch grains of various kinds in a series of fields, so that he can roughly estimate the amount of each starch present in the mixture, by careful comparison with mixtures of known percentage composition.

If the amount of starchy adulterant is considerable, evidence may be secured by determinations of starch by the diastase method and reducing matters by acid conversion.

Added Sugar.—Any appreciable amount of added cane sugar is shown by the sweet taste. The amount of cane sugar may be determined by means of the polariscope, as described on page 399.

An abnormally low ash is indicative of the addition of starch or sugar or both.

Foreign Fat.—Certain manufacturers have found it profitable to remove a portion of the cocoa butter from chocolate and substitute for it a cheaper fat, such as cocoanut oil, tallow or even paraffine. Such adulteration is detected by determination of the physical and chemical constants of the fat obtained by extraction with ether.

Dyes and Pigments, such as Bismark brown and Venetian red, have been employed to hide the presence of diluents. They are detected by dyeing tests, and by examination of the ash.

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CHAPTER XII.

SPICES.

THESE aromatic vegetable substances are classed as condiments, and depend for their use on the pungency which they possess in giving flavor or relish to food. As such seasoning or zest-giving substances, they are of considerable importance dietetically, but from the fact that they are used in comparatively insignificant amount, the determination of their chemical composition or actual value as nutrients per se is of little importance to the food economist.

Spices are, however, of chief interest to the public analyst, because of all food materials they constitute from their nature a class more susceptible than others to fraudulent adulteration of the most skilled variety.

In many cases not only the megascopic appearance and taste of the skillfully adulterated article are made to counterfeit the genuine spice, but even the microscopical appearance is intended to deceive, since it is the microscope that is most useful in the detection of adulteration, and in many cases in the determination of the approximate amount of the adulterants.

Indeed it is very rare that the microscope will fail to detect the presence of any foreign substance in spice, and hence its use is indispensable in the study of this class of foods by the analyst. Chemical methods, as a rule, while of secondary importance, are, however, very helpful, both as confirmatory of the microscopical research, and in some cases showing instances of adulteration not readily apparent with the microscope, such, for example, as in the case of exhausted spices, or those deprived of a whole or a part of their volatile oil. Sophistication of this kind is best shown by the ether extract.

General Methods of Proximate Analysis.—The following methods common to all the spices are for the most part those adopted provisionally by the A. O. A. C.* Methods peculiar to special spices will be treated

^{*} U. S. Dept. of Agric., Bur. of Chem., Bul. 65 and Bul. 107 (rev.).

under the discussion of the spice in question. For these determinations the spices should be powdered fine enough to pass through a 60-mesh sieve.

Determination of Moisture.—Richardson's Method.*—Two grams of the sample are weighed in a tared platinum dish and dried in an air-oven at 110° to a constant weight, which generally requires about twelve hours. The loss in weight includes the moisture and the volatile oil. The latter is determined from the ether extract, as described on page 410, and deducted from the total loss to obtain the moisture.

McGill † determines the moisture by exposure of a weighed portion of the sample *in vacuo* over perfectly colorless sulphuric acid. The spice gives up its moisture before the volatile oil comes off, and any appreciable amount of the volatile oil, when absorbed by the acid, causes the latter to be discolored, so that by carefully observing the beginning of the discoloration, and removing the sample, the loss due to moisture may be obtained by weighing at the proper stage. The abstraction of the moisture in this manner requires about twenty-four hours.

Determination of Ash.—Two grams of the spice are burned in a platinum dish heated to faint redness on a piece of asbestos paper by means of a Bunsen burner. The burning is best finished in a muffle furnace. If the ash contains an appreciable amount of carbon, it is exhausted on a filter with hot water, and the filter with the residue is burnt in the dish previously used. After adding the aqueous extract and a few drops of ammonium carbonate solution, the whole is evaporated to dryness and ignited at a faint red heat.

The Water-soluble Ash ‡ is found by boiling the total ash as above obtained with 50 cc. of water, and filtering on a tared Gooch crucible, the insoluble residue being washed with hot water, dried, ignited, and weighed. The insoluble ash, subtracted from the total, leaves the water-soluble ash.

Sand.—This is assumed to be the percentage of ash insoluble in hydrochloric acid. The ash from 2 grams of the substance, obtained as above described, is boiled with 25 cc. of 10% hydrochloric acid (specific gravity 1.050) for five minutes, the insoluble residue is collected on a tared Gooch crucible, thoroughly washed with hot water, and finally dried and weighed.

^{*} U. S. Dept. of Agric., Div. of Chem., Bul. 13, pt. 2, p. 165.

[†] Canada Dept. of Inland Rev. Bul. 73, p. 9.

[‡] U. S. Dept. of Agric., Bur. of Chem., Bul. 65, p. 55; Bul. 107 (rev.), p. 162.

Lime is determined from the ash as directed on page 303, having first separated the iron and phosphates.

The *sulphuric acid* due to calcium sulphate (added as an adulterant) is determined by precipitation with barium chloride of a very weak hydrochloric acid solution of the ash, the separated barium sulphate being washed, dried, ignited, and weighed.

Ether Extract.—Total, Volatile, and Non-volatile.*—Two grams of the air-dry, powdered substance are placed in some form of continuous extraction apparatus, such as Soxhlet's or Johnson's (pp. 63 and 68), and are subjected to extraction for sixteen hours with anhydrous, alcohol-free ether.† The ether solution is then transferred to a tared evaporating-dish, and allowed to evaporate spontaneously at the temperature of the room. After the disappearance of the ether, the evaporating-dish is placed in a desiccator over concentrated sulphuric acid and left over night, or for at least twelve hours, after which it is weighed, the residue in the dish being regarded as the total ether extract.

The dish and its contents are then subjected to a heat of about 100° C. for several hours, taking a long time to bring the temperature up to that point so as to avoid oxidation of the oil. Finally heat at 110° C. till the weight is constant. The final residue is the non-volatile, and the loss in weight the volatile ether extract.

Alcohol Extract.—Method of Winton, Ogden, and Mitchell.‡—Two grams of the powdered sample are placed in a 100-cc. graduated flask, which is filled to the mark with 95% alcohol. The flask is stoppered and shaken at half-hour intervals during eight hours, after which it is allowed to stand for sixteen additional hours without shaking, and the contents poured upon a dry filter. Of the filtrate, 50 cc. are evaporated to dryness in a tared platinum dish on the water-bath, and heated at 110° C. in an air-oven to constant weight. This method, while only approximate, is so much simpler than the tedious operation of continuous extraction, considering the long time required, that it is regarded as preferable for ordinary work, and, unless great care is taken, is nearly as accurate.

Determination of Nitrogen.—This, in spices other than pepper, is best done by means of the Gunning or Kjeldahl method (p. 69).

^{*} Richardson, U. S. Dept. of Agric., Div. of Chem., Bul. 13, p. 165.

[†] Petroleum ether may be used, yielding results which differ but slightly from those obtained with ethyl ether. As the latter has been used in the analyses of a large number of samples of spices, if these analyses are to be taken for standards of comparison it is evident that the same solvent should be used.

[‡] U. S. Dept. of Agric., Bur. of Chem., Bul. 65, p. 56; Bul. 107 (rev.), p. 163.

Determination of Starch.—In spices like white pepper, ginger, and nutmeg that normally contain a high content of starch and very little other copper-reducing matter, the direct acid conversion process of starch determination is satisfactory.

In spices normally free from starch, such as cloves, mustard, and cayenne, where a starch determination indicates the amount of a foreign starch present as an adulterant, it is safer to use the diastase process.

Four grams of the powdered sample are extracted on a filter-paper (fine enough to retain all starch particles) first with five successive portions of 10 cc. of ether, then with 150 cc. of 10% alcohol. Owing to difficulty of filtering in the case of cassia and cinnamon, Winton recommends that all washing in the determination of starch in these substances be omitted. The residue is washed from the filter-paper by means of a stream of water into a 500-cc. flask, if the direct acid conversion method is used, using 200 cc. of water; 20 cc. of hydrochloric acid (specific gravity 1.125) are added, and the method from this point on followed, as detailed on page 283.

If the starch is to be determined by the diastase method, wash the residue from the filter-paper into a beaker with 100 cc. of water, and proceed as on page 283.

Determine the dextrose in either case by the Defren or Allihn methods or volumetrically, and convert dextrose to starch by the factor 0.9.

Determination of Crude Fiber. — Two grams of the substance are extracted with ordinary ether (or the residue left from the determination of the ether extract may be taken) and subjected to the regular method for determining crude fiber, by boiling successively with acid and alkali (page 277).

McGill recommends the use of the centrifuge in separating the crude fiber, after boiling with the alkaline solution.

Determination of Volatile Oil.—Method of Girard and Dupré.*—
The spice is mixed with water and subjected to distillation, receiving the distillate in a graduated cylinder. The volume occupied by the essential oil (which is immiscible with water) can be thus read off and its content roughly determined. If the volatile oil is slightly soluble in water, separate out the water layer, having first read the volume of the oil layer, and extract the aqueous solution with petroleum ether. Evaporate the petroleum ether extract to dryness at room temperature

^{*} Analyse des Matieres Alimentaires, 2nd ed., p. 787.

in a tared dish, and add the volume due to the weight of the residue to the volume read off in the graduate.

Microscopical Examination of Powdered Spices.—As a rule few microscopical reagents are necessary in the routine examination of powdered spices for adulteration, unless a more careful study of the structure than is necessary to prove the presence of adulterants is desirable. The simple water-mounted specimen is usually sufficient to show the purity or otherwise of the sample. If in doubt as to the presence of starch in small quantities, iodine in potassium iodide should be applied to the specimen, well rubbed out under the cover-glass.

The tissues may be cleared by adding to the water mount a small drop of 5% sodium hydroxide, or by soaking a portion of the spice for a day in chloral hydrate solution. A valuable means of clearing dense tissues is to boil about 2 grams of the material successively with dilute acid and alkali as in the crude fiber process (p. 277), decanting (not filtering) the solution after each boiling.

The presence of occasional traces of a foreign substance, when viewed under the microscope, is hardly sufficient to condemn the sample as adulterated, since such traces are apt to be accidental.

Composition of Miscellaneous Spice Adulterants.—The chemical analyses of various spice adulterants commonly met with are given on page 413.

CLOVES.

Nature and Composition.—Cloves are the dried, undeveloped flowers of the clove tree (Caryophyllus aromaticus or Eugenia caryophyllata), which belongs to the myrtle family (Myrtacea). The tree is an evergreen, from twenty to forty feet in height, cultivated extensively in Brazil, Ceylon, India, Mauritius, the West Indies, and Zanzibar. Its leaves are from 7.5 to 13 mm. long, and its flowers, of a purplish color, grow in clusters. The green buds in the process of growth change to a reddish color, at which stage they are removed from the tree, spread out in the sun, and allowed to dry, the color changing to a deep brown. Each whole clove consists of a hard, cylindrical calyx tube, having at the top four branching sepals, surrounding a ball-shaped casing, which consists of the tightly overlapping petals, and within which are the stamens and pistil of the flower. In taste the clove possesses a strong and peculiar pungency. One of its most valuable ingredients is the volatile clove oil. This is composed largely of eugenol $(C_{10}H_{12}O_2)$, which forms 70 to

COMPOSITION OF SPICE ADULTERANTS.

			Ash.		Ether I	Extract.	act.
	Moisture.	Total.	Soluble in Water.	Insoluble in HCL	Volatile.	Non-vola- tile.	Alcohol Extract.
English-walnut shells * Brazil-nut shells * Almond shells * Cocoanut shells * Date stones * Spruce sawdust * Oak sawdust * Linseed meal * Cocoa shells * Red sandalwood * Ground olive stones † Buckwheat hulls.	7.69 9.08 7.80 7.36 8.24 8.77 5.73 8.71 10.44 4.42 9.50 7.63	1.40 1.59 2.86 0.54 1.24 0.23 1.22 5.72 8.40 0.70 0.88 1.84	0.77 1.06 2.39 0.50 0.76 0.16 0.32 1.74 4.66 0.28 0.24	0.00 0.17 0.05 0.00 0.04 0.00 0.02 0.55 0.83 0.07 0.44	0.12 0.07 0.16 0.00 0.36 0.07 0.04 1.00 1.21 0.06	0.55 0.57 0.64 0.25 8.38 0.77 0.84 6.58 2.99 11.47 0.24	1.84 1.01 5.16 1.12 16.72 1.50 6.25 9.46 4.77 19.37
	Reducing Matters by Acid Con- version.	Starch by Diastase Method.	Crude Piber.	Nitrogen X 6.25.	Oxygen Absorbed by Aqueous Extract.	Ouercitan- nic Acid Equivalent.	Total Nitro- gen.
English-walnut shells * Brazil-nut shells * Almond shells * Cocoanut shells * Date stones * Spruce sawdust * Oak sawdust * Linseed meal * Cocoa shells * Red sandalwood * Ground olive stones † Buckwheat hulls.	19.30 12.96 22.72 20.88 20.88 15.48 17.10 21.15 8.68 6.79	1.01 0.73 0.84 0.73 2.19 1.13 1.68 14.06 3.15 1.12 1.73 1.46	56.58 50.98 49.89 56.19 5.72 64.03 47.79 8.30 14.12 52.30 57.46 43.76	1.69 4.19 1.75 1.13 5.31 0.56 1.63 31.81 16.19 3.06 1.06 3.06	0.53 0.33 0.40 0.47 0.61 0.30 3.13 1.00 1.26	2.08 1.30 1.56 1.82 2.34 1.17 12.22 3.90 4.94 2.29	0.27 0.67 0.28 0.18 0.85 0.09 0.26 5.09 2.59 0.49

75 per cent of the oil, and a sesquiterpene known as caryophyllene. There are also in cloves a notable amount of fixed oil and resin, and also a peculiar form of tannin.

Very few complete analyses of cloves are on record. Richardson ‡ seems to have been the earliest worker in the field to give anything at all satisfactory in the way of a number of determinations of value.

The following are maximum and minimum figures from the tabulated results of Richardson's analyses:

^{*} Winton, Ogden, and Mitchell, Conn. Exp. Sta. An. Rep., 1898, p. 210.

[†] Doolittle, Mich. Dairy and Food Dept. Bul. 94, 1903, p. 18.

[‡] U. S. Dept. of Agric., Div. of Chem., Bul. 13.

	Water.	Ash.	Volatile Oil.	Fixed Oil and Resin.	Crude Fiber.	Albumin- oids.	Nitrogen.	Oxygen Equivalent.	As Querci- tannic Acid.
Whole cloves (7 samples): Maximum	10.67 2.90 10.18	5.50	18.89 10.23 4.40	7.12 4.03	6.18 13.58	4.73 5.78	.92	3.00 5.96	22.13 11.70 23.24
Maximum			13.93 3.94	7-44 4-02	13.80 9.38	6.48 4.20	1.04 .70		24.18

McGill* gives tables of analyses of pure and adulterated samples of cloves. Analyses of upwards of twenty samples of genuine cloves, both whole and ground, from these tables show the following maximum and minimum figures:

	Maximum.	Minimum.
Moisture	11.80	5.05
Volatile oil	19.63	9.24
Total volatile matter	30.68	16.25
Fixed oil.	10.23	0.94
Total extraction	31.40	22.23
Ash	7.00	5.03

McGill also made analyses of whole cloves of several varieties, the following table being a summary of his results:

		No. of Analyses.	Moisture.	Total Volatile Matter.	Volatile Oil.	Total Extract- ive Matter.	Fixed Oil.
Penang cloves:	Maximum Minimum	8	7-4 5.0	24.3 20.7	17.2	28.2 24.4	12.0
Amboyna cloves:	Mean Maximum Minimum	8	6.2 6.7 5.5	22.4 25.9 23.5	16.2 19.2 18.0	27.0 29.2 26.5	10.8 10.0 8.2
Zanzibar cloves:	Mcan	13	6. I 6. 7	24.6 23.6 18.6	18.5 18.3	27.5 28.1	9.0 10.7 8.0
	Mean	••••	4.1 5-7	21.7	16.0	21.3	9.6

Maximum and minimum figures of thirteen samples of unadulterated cloves, as purchased from retail dealers in Connecticut and analyzed by Winton and Mitchell,† are as follows:

^{*} Canada Inland Rev. Dept. Bul. 73.

[†] Conn. Exp. Sta. Rep., 1898, pp. 176-177

	Maximum.	Minimum.
Ash, total Ether extract, volatile. non-volatile	7.92 18.25 7.19	5.99 11.03 4.87

Winton, Ogden, and Mitchell * give more complete analyses of eight samples of whole cloves of known purity, representing Penang, Amboyna, and Zanzibar varieties, and two samples of clove stems, as follows:

		Ash.		Ether I	Extract.	Alcohol	
Moisture.	Total.	Soluble in Water.	Insoluble in HCl.	Volatile.	Non- volatile.	Extract.	
8.26 7.03 7.81 8.74	6.22 5.28 5.92 7-99	3.75 3.25 3.58 4.26	0.13 0.00 0.06 0.60	20.53 17.82 19.18 5.00	6.67 6.24 6.49 3.83	15.58 13.99 14.87 6.79	
Reducing Matters by Acid Conver- sion, as Starch.		Crude Fiber.	Nitrogen, ×6.25.	Oxygen Absorbed by Aque- ous Ex- tract.	Querci- tannic Acid.	Total Nitrogen.	
9.63 8.19 8.99 14.13	3.15 2.08 2.74 2.17	9.02 7.06 8.10 18.71	7.06 5.88 6.18 5.88	2.63 2.08 2.33 2.40	20.54 16.25 18.19 18.79	1.13 0.94 0.99 0.94	
	8.26 7.03 7.81 8.74 Reducing Matters by Acid Conversion, as Starch. 9.63 8.19 8.99	Reducing Matters by Acid Conversion, as Starch. 9.63 8.19 8.26 6.22 7.93 5.28 7.99 Reducing Matters by Acid Conversion, as Starch.	Moisture. Total. Soluble in Water. 8.26 6.22 3.75 7.03 5.28 3.25 7.81 5.92 3.58 8.74 7.99 4.26 Reducing Matters by Acid Conversion, as Starch. 9.63 3.15 9.02 8.19 2.08 7.06 8.99 2.74 8.10	Moisture. Total. Soluble in Water. Insoluble in HCl. 8.26 6.22 3.75 0.13 7.03 5.28 3.25 0.00 7.81 5.92 3.58 0.06 8.74 7.99 4.26 0.60 Reducing Matters by Acid Conversion, as Starch. 9.63 3.15 Crude Fiber. Nitrogen, X6.25. 9.63 3.15 9.02 7.06 8.19 2.08 7.06 5.88 8.99 2.74 8.10 6.18	Moisture. Total. Soluble in Water. Insoluble in HCl. Volatile. 8.26 6.22 3.75 0.13 20.53 7.03 5.28 3.25 0.00 17.82 7.81 5.92 3.58 0.06 19.18 8.74 7.99 4.26 0.60 5.00 Reducing Matters by Acid Conversion, as Starch Crude Fiber. Nitrogen, X6.25. Oxygen Absorbed by Aqueous Extract. 9.63 3.15 9.02 7.06 2.63 8.19 2.08 7.06 5.88 2.08 8.99 2.74 8.10 6.18 2.33	Moisture. Total. Soluble in Water. Insoluble in HCl. Volatile. Non-volatile. 8.26 6.22 3.75 0.13 20.53 6.67 7.03 5.28 3.25 0.00 17.82 6.24 7.81 5.92 3.58 0.06 19.18 6.49 8.74 7.99 4.26 0.60 5.00 3.83 Reducing Matters by Acid Conversion, as Starch Crude Fiber. Nitrogen, X6.25. Oxygen Absorbed by Aqueous Extract. Quercitannic ous Extract. 9.63 3.15 9.02 7.06 2.63 20.54 8.19 2.08 7.06 5.88 2.08 16.25 8.99 2.74 8.10 6.18 2.33 18.19	

The Tannin Equivalent in Cloves.—The amount of tannin in cloves was shown by Ellis to be so constant as to be of valuable assistance as a guide to their purity. The actual determination of tannin is, however, a long and difficult proceeding, and Richardson† has pointed out that it is not necessary, but that simply using the first part of the Lowenthal tannin process, and noting the "oxygen absorbed" as expressed by the oxidizing power of permanganate of potash on the material after extraction with ether, is quite as useful as determining the tannin, and is in effect proportional to the tannin present. The result is sometimes expressed as in Richardson's figures above, as the oxygen equivalent, or as quercitannic acid.

Determination of Tannin Equivalent.‡—Reagents: Indigo Solution.— Six grams of the indigo salt § are dissolved in 500 cc. of water by heat-

^{*} Conn. Exp. Sta. Rep., 1898, pp. 206, 207.

[†] U. S. Dept. of Agric., Div. of Chem., Bul. 13, p. 167.

[‡] U. S. Dept. of Agric., Bur. of Chem., Bul. 65, p. 60; Bul. 107 rev., p. 164.

[§] The quality of the indigo used is of great importance since with inferior brands it is

ing. After cooling, 50 cc. of concentrated sulphuric acid are added, the solution made up to a liter and filtered.

Standard Permanganate Solution.—Dissolve 1.333 grams of pure potassium permanganate in a liter of water. This should be standardized by titrating against 10 cc. of tenth-normal oxalic acid (6.3 grams pure crystallized oxalic acid in 1,000 cc.), diluted to 500 cc. with water, heated to 60° C., and mixed with 20 cc. of dilute sulphuric acid (1:3 by volume). The permanganate solution is added slowly, stirring constantly, till a pink color appears.

Two grams of the material are extracted for twenty hours with pure anhydrous ether. The residue is boiled for two hours with 300 cc. of water, cooled, made up to 500 cc., and filtered.

Twenty-five cc. of the filtrate are pipetted into a 1200-cc. flask, 750 cc. of distilled water are added and 20 cc. of indigo solution.

The standard permanganate solution is then run in from a burette a drop at a time with constant shaking, until a bright golden yellow color appears, which indicates the end-point. Note the number of cubic centimeters required, represented by (a).

In a similar manner determine the number of cubic centimeters of standard permanganate solution consumed by 20 cc. of the indigo solution alone, represented by (b), and subtract this from (a).

The oxygen equivalent, or, as it is sometimes called, the "oxygen absorbed," is calculated from the equivalent in tenth-normal oxalic acid of the number of cubic centimeters of standard permanganate represented by a-b. 10 cc. of tenth-normal oxalic acid are equivalent to 0.008 gram of oxygen absorbed, or 0.0623 gram of quercitannic acid.

Microscopical Examination of Cloves.—Unless the finely powdered, water-mounted sample is well rubbed out under the cover-glass, many of the masses of cellular tissue will be too dense to recognize. With a little care, however, it is possible to make a very satisfactory water mount, though by soaking for twenty-four hours in chloral hydrate solution the more opaque masses are rendered very translucent.

Fig. 81, from Moeller, shows some of the characteristics of powdered cloves. The outer skin of the calyx tube is shown at (1) with its polygonal cells and large oil spaces showing through them; (2) shows the epidermis of the outer part of the lobes or wings of the calyx, with stomata

impossible to get a sharp end-point. The indigo solution should be made from the very best variety of sulphindigotate, which may be obtained from Grueber & Co., of Leipzig, or Gehe & Co., of Dresden, under the name of carminium caruleum.

surrounded by irregularly shaped cells; (3) represents the epidermis of the petals, with crystals of calcium oxalate; a cross-section of the epidermis of the calyx is shown at (4); (5) shows the parenchyma, with calcium oxalate crystals and with one of the slender spiral ducts; (6) and (7) represent in cross-section and longitudinal section respectively the parenchyma of the middle layers of the ovary, one of the rounded, triangular pollen grains being shown at (12).

Fig. 81.—Powdered Cloves under the Microscope. X125. (After Moeller.)

Characteristics of clove stems, which are frequently used as adulterants of cloves, are found in (8), (9), (10), and (11). Stone cells of the outer skin and the inner portion of the clove stem are shown at (8) and (9) respectively; (10) shows one of the vascular ducts, and (11) two of the bast fibers. Both the vascular ducts and the stone cells are very characteristic of clove stems. Pure cloves have no stone cells and comparatively few bast fibers. Stems under the microscope show a large number of bast fibers and frequent stone cells, the latter being of a distinctly yellow color.

A plain water-mounted slide rarely shows all the structural details depicted in Fig. 81, but is nearly always sufficiently characteristic to

prove the purity of the sample. Fig. 220, Pl. XXV, shows the actual appearance of powdered cloves, mounted in water and examined under a magnification of 130. The general appearance of the cellular tissue is that of a loose, spongy mass filled with brown, granular material. Throughout the masses of tissue are to be seen small oil globules.

Cloves have no starch whatever. Aside from the stems, cloves are sometimes adulterated with clove fruit or "mother cloves," which have a small amount of a sago-like starch, and also contain some stone cells.

Adulteration of Cloves.—The U. S. standard for pure cloves is as follows: Volatile ether extract not less than 10%; quercitannic acid, calculated from the total oxygen absorbed by the aqueous extract, should not be less than 12%; total ash should not exceed 8%; ash insoluble in hydrochloric acid should not exceed 0.5%, and crude fiber should not be more than 10%.

Clove Stems are very frequent adulterants of cloves and possess some slight pungency. They are commonly identified under the microscope by the large number of bast fibers and stone cells, and should not be found in pure cloves in excess of 5%.

Allspice, being considerably cheaper than cloves, is sometimes used as an adulterant. It is readily recognized by the characteristics described on page 422.

Other Adulterants commonly found are cereal starches (especially corn and wheat) and ginger (for the most part "exhausted"). Besides the above, pea starch, rice, turmeric, charcoal, sand, pepper, ground fruit stones, and sawdust have been found in samples of cloves examined in Massachusetts.

Exhausted Cloves, both whole and in powdered form, are not infrequently found on the market. These have been deprived of a portion of the volatile oil, and are much less pungent than the pure article, so that the difference in taste between the two varieties is quite marked. It is, however, rare that powdered cloves are sold consisting entirely of the exhausted variety, the more common practice being to mix from 10 to 25 per cent of exhausted cloves with the pure powder, so that the sophistication is less apparent.

A determination of the volatile oil is the only reliable means of showing whether or not the material has been wholly or in part exhausted, though Villier and Collin claim that under the microscope an exhausted sample of cloves shows the oil glands to be nearly empty, or to inclose much smaller droplets of oil than the pure variety.

With the exception of exhausted cloves, the presence of nearly every foreign ingredient is best and most quickly shown by the use of the microscope, though much information as to the purity of the sample can be gained by the ether extract, the percentage of ash, and of crude fiber.*

Cocoanut Shells.—Figs. 226 and 227, Pl. XXVII, show samples of cloves adulterated with ground cocoanut shells. The long, spindle-shaped, yellow-brown and deeply furrowed stone cells of the adulterant with their thick walls and central branching pores are unmistakable. The dark-brown contents of the cells turn reddish brown when treated with potassium hydroxide. The anatomy of the cocoanut, including the shell, has been carefully studied by Winton.†

Fig. 82, after Winton, shows elements of powdered cocoanut shell under the microscope. st are the dark, elongated, yellow, porous stone

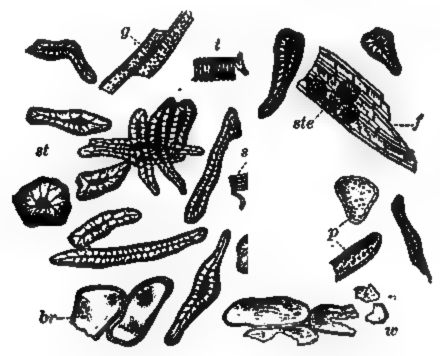


Fig. 82.—Cocoanut-shell Powder. st, dark-yellow stone cells with brown contents; t, reticulated trachea; sp, spiral trachea; g, pitted trachea; w, colorless, and br, brown, parenchyma of mesocarp; f, bast fibres, with stegmata (ste). ×160. (After Winton.)

cells with their brown contents, these stone cells being the most distinctive characteristic of the ground shells. t, sp, and g are the various forms of trachea; w and br are respectively colorless and brown parenchyma of the mesocarp or outer coat, portions of which always adhere to the nutshell and are ground with it.

^{*} Note especially the sharp distinction between these values in the case of pure cloves and of clove stems in Richardson's table.

[†] The Anatomy of the Fruit of the Cocoanut. Conn. Exp. Sta. Rep., 1901, p. 208.

Fig. 264, Pl. XXXVI, shows a photomicrograph of powdered cocoanut shells, mounted in gelatin. The long, spindle-shaped stone cells are especially apparent,

Ground cocoanut shells have been used in various spices besides cloves, especially allspice and pepper. In the following tabulated results of analyses by Winton, Ogden, and Mitchell * are shown the wide deviation between the chemical constants of cocoanut shells and several of the spices in which they appear as adulterants.

	Black Pepper.	Cloves.	Allspice.	Nutmeg.	Cocoanut Shells.
Water	11.96	7.8r	9.78	3.63	7.36
Total ash	4.76	5.92	4.47	2.28	0.54
Ash soluble in water	2.54	3.58	2.47	0.86	0.50
Ash insoluble in hydrochloric acid	0.47	0.06	0.03	0.00	0.00
Volatile ether extract	1.14	19.18	4.05	3.02	0.00
Non-volatile ether extract	8.42	6.49	5.84	36.70	0.25
Alcohol extract	9.62	14.87	11.79	10.77	1.12
Reducing matters, as starch, acid conversion	38.63	8.99	18.03	25.56	20.88
Starch by diastase method	34.15	2.74	3.04	23.72	0.73
Crude fiber	13.06	8.10	22.30	2.51	56.10
Total nitrogen	2.26	0.99	0.02	1.08	0.18
Oxygen absorbed by aqueous extract		2.33	1.24	l	0.23
Quercitannic acid equivalent		18.19	9.7i		1.83

ALLSPICE, OR PIMENTO.

Nature and Composition.—Allspice is the dried fruit of the Eugenia pimenta, an evergreen tree belonging to the same family (Myrtacea) as the clove. It is indigenous to the West Indies, and is especially cultivated in Jamaica.

The allspice berry is grayish or reddish brown in color, and is hard and globular, measuring from 4 to 8 mm. in diameter, being surmounted by a short style. This is imbedded in a depression, and around it are the four lobes of the calyx, or the scars left by them after they have fallen off. The berry has a wrinkled, ligneous pericarp, with many small excrescences filled with essential oil. The pericarp is easily broken between the fingers, showing the berry to be formed of two cells with a single, brown, kidney-shaped seed in each, covered with a thin, outer coating, inclosing an embryo rolled up in a spiral.

The berries are gathered when they have attained their largest size, but before becoming fully ripe. If allowed to mature beyond this stage, some of the aroma is lost.

^{*} Conn. Ag. Exp. Sta. Rep., 1901, p. 225.

Though considerably less pungent than other spices, allspice possesses an aroma not unlike cloves and cassia. In chemical composition it most resembles cloves, containing both volatile oil and tannin; but, unlike cloves, it contains much starch, the starch being contained in the seeds. The volatile oil of allspice is very similar to clove oil. It is slightly lævorotary, and is composed of eugenol and a sesquiterpene not determined. It is present in allspice to the extent of 3 to 4.5 per cent. The boiling-point of the oil is 255° C.

Authoritative full analyses of allspice are even more meager than of cloves. Analyses of one sample of whole allspice and five samples of the ground spice, made by Richardson,* are thus summarized:

	Water.	Ash.	Volatile Oil.	Fixed Oil.	Undeter- mined.	Crude Fiber.	Albumin- oids.	Nitrogen.	Tannin Equivalent.	Oxygen. Required.
Whole	6.19 8.82 5.51	5.53 3.45	3.3 ² 2.07	6.15 6.92 3-77	59.28 58.24 56.86	18.98	5-42	.70 .87 .64	10.97 12.74 8.27	3.36

Seventeen samples of unadulterated allspice, as sold on the Connecticut market, were analyzed by Winton and Mitchell,† with maximum and minimum results as follows:

Ash.	Maximum.	Minimum.
Total. Insoluble in hydrochloric acid (sand) Ether extract, volatile. Ether extract, non-volatile	3.50	4-34 -40 1-34 3-78

Three samples of pure whole allspice were more fully analyzed by Winton, Mitchell, and Ogden with the results given on page 422.

The Tannin Equivalent in Allspice.—Tannin is present in allspice, though to a less extent than in cloves. The exact amount present is rarely determined, but rather the "oxygen equivalent," or quercitannic acid, as explained on page 415, the determination being carried out as there detailed.

^{*} U. S. Dept. of Agric., Div. of Chem., Bul. 13, p. 229.

[†] An. Rep. Conn. Exp. Sta., 1898, pp. 178, 179.

¹ Ibid., pp. 208, 209.

			Ash.		Ether I	Alcohol	
	Moisture.	Total.	Soluble in Water.	Insoluble in HCl.	Volatile.	Non- volatile.	Extract.
Maximum	10.14 9.45 9.78	4.76 4.15 4.47	2.69 2.29 2.47	o.o6 o.oo o.o3	5.21 3.38 4.05	7-72 4-35 5-84	14.27 7.39 11.79
	Reducing Matters by Acid Conver- sion, as Starch.	Starch by Diastase.	Crude Fiber.	Nitrogen, X 6.25.	Oxygen Absorbed by Aque- ous Ex- tract.	Querci- tannic Acid.	Total Nitrogen.
Maximum	20.65 16.56 18.03	3.76 1.82 3.04	23.98 20:46 22.39	6.37 5.19 5.75	I.59 I.03 I.24	12.48 8.06 9.71	1.02 0.83 0.92

Microscopical Examination of Powdered Allspice.—By soaking the powder twenty-four hours or more in chloral hydrate, many of the harder portions are rendered much more transparent than would otherwise be possible. Fig. 83, after Moeller, shows the microscopical structure of various elements that go to make up allspice powder.

The epidermis, or outer layer of the berry, is shown at (1a) in cross-section, and in plan view at (2) with its small cells. Just beneath the outer coat are the large oil spaces (1b) and still further below the stone-cells (1c). The fruit parenchyma (3) has vascular tissues running through it. (4) and (5) are the inner epidermis and stone cells of the dividing partitions between the seeds. Small hairs connected with the outer epidermis are shown at (6). (7) and (8) show in cross-section a portion of the seed-shell and inclosed seed or embryo, with the starch (8a) and the colored lumps of gum or resin (8b) of a port-wine color. These colored cells exist in the seed coating, and, although only one is here shown, constitute a very important and striking characteristic of allspice. (9) represents the spongy parenchyma of the seed shell, and (10) shows its epidermis. In the parenchyma of the fruit and of the partitions between the cells are seen, but not always plainly, minute crystals of calcium oxalate (see (4) and (5)).

These details so closely drawn by Moeller are idealized, but serve well to indicate what should be looked for. In practice the water-mounted specimen shows all the characteristics necessary to identify pure allspice, and most if not all its adulterants. In fact pimento is one of the easiest spices to identify under the microscope, by reason of its striking characteristics.

Three distinctive features are especially typical, viz.: First, the starch grains, which are very uniform in size, measuring about 0.008 mm. in diameter, being nearly circular as a rule, and often arranged in groups not unlike masses of buckwheat starch. Ordinarily these masses contain fewer granules than do those of buckwheat. The granules are



b

20.00

Fig. 83.—Powdered Allspice under the Microscope. X125. (After Moeller.)

smaller and more inclined to the circular than to the polygonal form, while in many cases they have distinct central hila. The starch grains are very numerous and are found in nearly every field. See Fig. 195, Pl. XIX.

A second distinctive feature of allspice is the stone cells, of which there are many. These are more often colorless, and in most cases very large and plainly marked. They are sometimes seen singly and at other times grouped together. Frequently they are attached to pieces of brown parenchyma.

The third and most characteristic feature of allspice powder under the microscope is the striking appearance of the lumps of gum or resin, which are of a more or less deep port-wine or amber color and are contained in the middle layers of the seed coat. These cells are very striking, occurring sometimes in isolated bits, and in other cases in aggregations of from 2 to 4 or even 6 to 8 cells. These resinous lumps appear plainly in Fig. 194, Pl. XIX. Droplets of oil are occasionally seen, but not in profusion. As a rule the oil is forced out of its large containing cells and into the surrounding tissue by the process of drying.

Adulteration of Allspice.—According to the U. S. standard for allspice, quercitannic acid should not be less than 8%, total ash not more than 6%, ash insoluble in hydrochloric acid not more than 0.5%, crude fiber not more than 25%. The most common adulterants found in powdered allspice are cocoanut shells and the cereal starches. Besides these the writer has found in Massachusetts, peas, pea hulls, exhausted ginger, cayenne, olive stones, pepper, and turmeric. To this list may be added clove stems, which are on record as a not uncommon adulterant in some localities. All of these are to be readily recognized by a careful microscopical examination.

CASSIA AND CINNAMON.

Nature and Composition. — The terms cassia and cinnamon are interchangeable in commerce, though, strictly speaking, they represent two separate and distinct species of the genus Cinnamomum, belonging to the laurel family (Lauracea). True cinnamon is the bark of Cinnamomum zeylanicum, a tree from 20 to 30 feet high, having horizontal or drooping branches, and native to the island of Ceylon, but cultivated also in some parts of tropical Asia, in Sumatra, and in Java. The entire yield of pure Ceylon cinnamon is extremely small, and but little of it is found in this country. It is the very thin, inner bark of the tree, and is of a pale, yellowish-brown color, being found on the market in long, cylindrical, quill-like rolls or pieces, the smaller rolls being inclosed in the larger. The outer surface is marked by round dark spots, corresponding to points of insertion of the leaves, and it is also furrowed lengthwise by somewhat wavy, light-colored lines. The inner surface of the bark is darker colored, and has no lines. In thickness the bark varies. from 1.5 to 3 mm. Both the inner and outer coatings of the bark of Ceylon cinnamon are usually removed in the process of preparation, so

that it is of a much cleaner and more even texture than the cassia bark, which is thicker and heavier by reason of the outer cork layer usually left on it.

The cheaper and more common cassia is the bark of the Cinnamomum cassia, which comes from China, Indo-China, and India. It is of a darker color than that of cinnamon, of coarser texture, and as a rule about four times as thick. Most varieties of cassia bark are less tightly rolled than cinnamon, and are not arranged one within the other in layers. The outer surface is marked by elliptical spots left by the leaves, and by small, dark-brown, wart-like protuberances. Cassia does not have the wavy, light-colored lines found in the cinnamon. Both cinnamon and cassia barks are very aromatic in taste, somewhat astringent, and slightly sweet.

Cassia buds are the dry flower buds of China cassia, and are found in the market both in whole and in powdered form. Powdered cassia often consists of a mixture of several varieties of bark, while the cheaper grades sometimes contain an admixture of the ground buds.

The best grade of cassia is that from Saigon, a much cheaper, from Batavia, while the cheapest is the China cassia.

The odor of cassia and cinnamon bark is due to the volatile oil, of which from 1 to 2 per cent is usually found. Cassia and cinnamon oil greatly resemble each other, the principal constituent in either case being cinnamic aldehyde, C₆H₅CH: CH.CHO. Besides this, one or more esters of acetic acid are present. Both oils are very pungent and intensely sweet.

Starch is present in cassia to the extent of from 16 to 30 per cent. A very small amount of tannin is found, as well as cinnamic acid and mucilaginous matters. Cassia buds are somewhat similar in composition to the bark. They have, however, less starch and crude fiber, and higher contents of volatile oil and nitrogen than the bark.

Richardson* has made analyses of a few samples of pure whole cinnamon and cassia, from which the following are taken:

	Water.	Ash.	Volatile Oil.	Fixed Oil, etc.	Crude Fiber.	Albumin- oids.	Undeter- mined.	Nitrogen.
Ceylon cinnamon, 1	5.40 7.43 4.79 17.45 9.32	3.40 5.58 8.23	1.05 .82 3-59 3-51 -55	1.66 1.58 5.21 2.38 -74	33.08 25.63 8.60 26.29 14.33	3.80 7.00 4.55	51.28 56.84 65.23 65.33 48.65	.48 .62 1.12 .73

* U. S. Dept. of Agric., Div. of Chem., Bul. 13, p. 221.

Winton, Ogden, and Mitchell's * results of analyses of whole samples of cinnamon, cassia, and cassia buds are thus summarized:

	Moisture.		Ash.		Ether Extract.		
		Total.	Soluble in Water.	Insoluble in HCl.	Volatile.	Non- volatile	
Ceylon cinnamon (6 samples):							
Maximum	10.48	5-99	2.71	0.58	1.62	r.68	
Minimum	7-79	4.16	1.40	0.02	0.72	1.35	
Average	8.63	4.82	1.87	0.13	1.39	1.44	
Cassia bark (20 samples):			1				
Maximum	11.91	6.20	2.52	2.42	5.15	4.13	
Minimum	6.53	3.01	0.71	0.02	0.93	1.32	
Average	9.24	4-73	1.68	0.56	2.61	2.12	
Cassia buds (2 samples):				1			
Average	7-93	4.64	2.88	0.27	3.88	5.96	

	Alcohol Extract.	Reducing Matters by Acid Conversion, as Starch.	Crude Fiber.	Nitrogen, X 6.25.	Total Nitrogen.
Ceylon cinnamon (6 samples): Maximum	13.60	22.00	38.48	4.06	0.65
Minimum	9-97	16.65	34.38	3.25	0.52
Average	12.21	19.30	36.20	3.70	0.59
Maximum	16.74	32.04	28.80	5-44	0.87
Minimum	4-57	16.65	17.03	3.31	0.53
Average	8.29	23.32	22.96	4-34	0.69
Cassia buds (2 samples): Average	10.88	10.71	13.35	7-53	1.20

Structure of Powdered Cassia under the Microscope. — Fig. 84, from Moeller, shows various elements of cassia bark as veiwed microscopically. (1) shows in cross-section a portion of the cork and outer layer of the bark rind, with flat cells nearest the surface, having somewhat thick walls and reddish-brown contents, and, farther in, the cells s, with mucilaginous material.

The stone cells of the intermediate layer of bark are shown at (2). Here the tendency of the stone cells is to be thicker on one side than on the other, as is plainly shown. (3) represents the structure of the inner layer of the bark, showing bast fibers b cut across, and more of the so-called mucilaginous cells s of large size, which normally contain the ethereal or volatile oil. The starch granules (4) are contained in great abundance in the polygonal cells of the parenchyma of the intermediate

^{*} Twenty-second Annual Report Conn. Exp. Sta., 1898, pp. 204, 205.

and inner bark layers. (6) represents a fragment of a bast fiber, which is often shown in cassia powder with connecting parenchyma. The stone-cells of the cork are shown in plan view at (7). Very small, needle-like crystals of oxalate of calcium are occasionally to be seen if looked for carefully. They occur in the parenchyma cells of the inner and intermediate layers of the bark.

The microscopical structure of Ceylon cinnamon much resembles that of cassia. Cassia starch grains measure from 0.0132 to 0.0222 mm.,

1



Fig. 84.—Powdered Cassia under the Microscope. X125. (After Moeller.)

being considerably larger and more abundant that those of true cinnamon. As a rule the bast fibers of cassia are larger, but shorter, than those of cinnamon, and provided with thicker walls.

Figs. 203 and 204, Pl. XXI, show various phases of pure cassia bark as photographed from water-mounted specimens of the powder. Cassia starch somewhat resembles that of allspice, but it is not as a rule found in masses containing as many granules as does the allspice starch. Very commonly two or three of the starch granules are arranged together in

such a manner that at first sight they appear to form a single large granule, but on more careful examination are seen to be two- and three-lobed, consisting of several smaller grains. Stone cells, which are very abundant in the powdered cassia, do not happen to be included to any extent in the photographed fields. Cassia stone cells are generally more oblong than those of allspice, and are more often brown in color, while the allspice stone cells are generally colorless.

A distinctive feature of powdered cassia consists in the long, ambercolored wood fibers, some distributed in bundles, and others arranged singly. These are very clearly shown in Figs. 204 and 205.

Yellow patches of cellular tissue with starch grains interspersed among them are very abundant in the powder.

Adulteration of Cinnamon and Cassia.—The U. S. standards are as follows: Total ash not to exceed 8%; sand not to exceed 2%.

The commonest adulterants are cereal products and foreign bark. Besides these, the writer has found, in samples sold in Massachusetts, leguminous starches, pea hulls, nutshells, turmeric, pepper, olive stones, ginger, mustard, and sawdust. Much of the China cassia when imported contains an inexcusably large amount of dirt. In one sample Winton, Ogden, and Mitchell found over 15% of sand.

Ground Bark of the Common Trees, especially that of the elm, resembles in physical appearance ground cassia, and is to be looked for as an adulterant. Fig. 265, Pl. XXXVII, shows the appearance of ground elm bark. The fibers of cassia bark have starch granules as a rule interposed among them, while the foreign bark, usually of a much coarser texture, shows no starch connected with its structure.

Fig. 206, Pl. XXII, shows a water-mounted specimen of adulterated cassia powder, chosen from samples purchased in the Massachusetts market. Nothing but the adulterant (a foreign bark) shows in the field. The tissue is loose and considerably coarser than that of cassia bark.

PEPPER.

Nature and Composition.—Pepper is the dried berry of the pepper plant (*Piper nigrum*), a climbing shrub belonging to the family *Piperacea*, native to the East Indies, but cultivated in many tropical countries. The height of the pepper plant is from twelve to twenty feet. When the fruit begins to turn red, it is gathered and then dried, by which processit turns black and shrivels up, forming the black peppercorns of commerce. They are spherical single-seeded berries, about 5 mm. in diam-

eter, covered with a brownish-gray epicarp, and having on the under side the remains of a short stem. At the top of the berry is an indistinct trace of a style, and of a lobed stigma.

Varieties of black pepper are named from the localities in which they are grown or from which they are shipped, as Singapore, Lampong, Sumatra, Tellichery, Malabar, Acheen, Penang, Alleppi, Trang, Mangalore, etc.

White pepper is obtained by decorticating the fully ripened black peppercorns, or removing the dark skin. This is accomplished by macerating them in water to loosen the skin, which is then removed readily by drying and rubbing between the hands. White whole pepper grains are grayish white, and a trifle larger than the black pepper berries. They are nearly spherical in shape, and have a number of light-colored lines that, like meridians, run from top to bottom. The common varieties are Siam, Singapore and Penang, the latter being coated with lime.

The pungent taste of pepper is due in great part to its essential oil, a hydrocarbon of the formula $C_{10}H_{16}$, present in amounts varying from 0.5 to 1.7 per cent. Pepper oil contains phellandrene and a terpene.

Other important constituents of pepper are piperidine, and the crystalline base *piperin*, C₁₇H₁₉NO₃, insoluble in water, but soluble in ether, and in alcohol. Starch is present in pepper to a large extent.

Burcker gives the following average percentage composition of black and white pepper:

	Ash.	Cellulose.	Water.	Nitrogenous Matter.	Volatile Oil.	Fat, Piper- in, and Resin.	Starch and Dextrin.	Other Non- nitrogenous Matters.
Black pepper	4·57 1.80	12.45 6.08	12.50 13.56	11.98	1.36	6.85 7.11	42.90 56.04	7-39 3-35

Richardson's * analyses of three samples of whole black and two samples of whole white pepper, all pure, are as follows:

	Water.	Ash.	Volatile Oil.	Piperin and Resin.	Alcohol Extract.	Starch (Acid Con- version)
Black pepper: West coast	8.91 8.29	4.04	.70 1.69	7-29 7-72	6.06	36.52 37.50
Singapore White pepper: West coast Singapore	9.83 9.85 10.60	3.70 1.41 1.34	1.60 -57 1.26	7.15 7.24 7.76	5-74 2-57	37.30 40.61 43.10

^{*} U. S. Dept. of Agric., Bur. of Chem., Bul. 13, part 2, p. 206.

		Undeter- mined.	Crude Fiber.	Albumin- oids.	Total N×6.25.	Total N.
	West coast	23.28	10.23 10.02 10.02 7.73	7.69 10.38 10.00 9.31	9.81 12.60 12.08 11.48	1.57 2.02 1.93 1.83
White pepper:	West coast Singapore	23.28		1	1	

Richardson gives the following variations in the constituents of pure pepper:

	Black.	White.
Water	8.0 to 11.0	8.0 to 11.0
Ash	2.75 to 5.0	1.0 to 2.0
Volatile oil	.50 to 1.75	.50 to 1.75
Piperin and resin	7.0 to 8.0	7.0 to 8.0
Starch	32.0 to 38.0	40.0 to 44.0
Crude fiber.	8.0 to 11.0	4.11 to 8.0
Albuminoids	7.0 to 12.0	8.0 to 10.0

McGill's * analyses of six samples of whole black, and five samples of whole white pepper, all genuine, are thus summarized:

				Ash.			
	Moisture, etc., Lost at 100° C.	Soluble in Hot Water.	Insoluble in Water.	Total.	Insoluble in Hydro- chloric Acid.	Sand Expressed as Per Cent of Total Ash.	Alcohol Extract.
Black: Maximum	14.10	2.64	3.06	5.16	1.08	21	9.06
Minimum	10.62	2.07	1.46	3.98	.06	2	8.28
Mean	12.03	2.41	2.05	4-47	0.36	8	8.71
White: Maximum	13.00	0.72	3.04	3.65	0.88	42	8.92
Minimum	11.30	0.14	1.50	1.64	0.26	9	7.00
Mean	12.34	0.54	2.46	3.00	0.55	21	7.73

Winton, Ogden, and Mitchell's, and Winton and Bailey's † analyses of whole black pepper and whole white pepper, representing the leading varieties imported into the United States, also of pepper shells and long pepper, are summarized in the following table:

^{*} Canada Inl. Rev. Dept. Bul. 20, 1890.

[†] An. Rep. Conn. Exp. Sta., 1898, pp. 198-199; 1903, pp. 158-164.

·				Ash.		Ether Extract	xtract.		r, 0)			N :	Nitrogen.	gen.
	Number of Samples	Moisture.	TetoT	Soluble in Water.	Insoluble in HCl.	Volatile.	Non-volatile.	Alcohol Extract.	Reducing Matters b Acid Conversion as Starch.	Starch by Diastase.	Crude Piber.	Total Nitrogen, less in Ether Extract, ×6.25.	.latoT	In Non-volatile Ether Extract.
Whole Black Pepper:														
SingaporeAverage	v	12.00	3.49	2.10	0.12	90.1	7.73	8.89	42.80	38.51	11.04	12.71	2.30	0.31
	4	11.56	4.21	2.75	10.0	0.83	6.94	8.8	41.67	37.01	12.20	11.79	2.15	0.27
LampongAverage	4	11.46	6.05	2.37	1.06	1.22	8.57	9.41	39.66	35.08	12.37	10.90	2.08	0.33
Acheen A*	H	12.09	5.04	2.78	0.48	1.09	6.17	10.04	38.17	33.30	13.07	10.88	2.11	0.37
Acheen BAverage	3	12.41	5.65	2.91	1.08	1.49	8.59	9.30	37.11	32.17	14.80	11.79	2.24	0.35
Acheen C Average	v	11.78	5.99	3.06	1.05	1.64	6.56	10.15	31.97	25.76	17.19	12.32	2.35	0.38
All varietiesMaxinum	8	12.95	6.85	8.80	1.63	28.80	10.37	11.86	43.47	39.66	18.25	18.81	2.53	0.39
Minimum		10.63	8.09	1.75	0.00	99.0	6.86	8.31	28.15	22.05	10.75	10.50	£.03	0.87
Average		11.86	6.10	2.60	0.70	1.28	8.41	9.44	38.28	33.28	13.62	11.93	38.8	0.33
Whole White Pepper:														
DecorticatedAverage	01	12.89	90.1	0.47	100	0.56	7.24	7.83	64.85	63.16	9.6	11.03	3.08	0.32
SingaporeAverage	"	13.47	1.33	0.33	0.00	0.92	7.89	8.45	56.71	53.89	4.10	11.06	2 11	0.34
	8	13.63	1.47	0.38	0.10	69.0	6.64	7.36	59.01	56.18	3.52	10.69	700	0.39
Penang Average	8	13.68	2.84	0.65	0.15	92.0	6.33	7.32	57.17	54.01	3.80	10 88	2.01	0.27
All VarietiesMaximum	o.	14.47	2.96	080	0.30	0.95	7.94	8.55	86.79	63.60	4.85	11.19	2.13	0.34
Minimum		12.72	1.03	0.28	0.00	67.0	8.86	7.19	56.43	53.11	79.0	10.44	1.95	93.0
Average		13.47	1.77	0.47	0.10	0.73	16.9	2.66	59.17	26.47	8.14	10.89	2.04	0.30
Pepper ShellsMaximum	ĸ	10.66	16.11	3.20	4.70	90.1	4.97	6.30	21.69	15.30	32.15	14.19	2.36	0.15
Minimum		10.52	10.25	2.28	2.63	90.0	3.04	8.	11.43	2.30	23.27	12.31	2.12	6.00
Whole Long Pepper.	н	9.47	5.93	4.30	0.33	1.55	19.9	8.67	42.88	39.55	5.76	12.25	2.18	0.33
			-				_				_		-	

* Acheen pepper contains bollow kernels and shells varying in amount with the grade.

The following table summarizes the results of full analyses of pepper and pepper shells recently made by Doolittle:*

	N 6	N	Mois-		Ash.		Starch by
	No. of Samples.	No. of Varieties.	ture.	Total.	Insoluble in HCl.	Soluble in Water.	Diastase Method.
Black pepper:	45	12					
Maximum	٠		11.96	8.04	2.59	3.32	41.75
Minimum			8.00	3-43	0.05	1.65	25.00
Average	l	••••	9.54	4-99	0.58	2.49	36.69
White pepper:	25	9	' • '	' ' '		1	
Maximum			13.34	4.28	0.86	1.16	63.55
Minimum			8.04	0.86	0.05	0.12	63.55 48.88
Average			9.87	1.69	0.19	0.34	54-37
Long pepper:	3		, ,	1	1		
Maximum			10.13	14.39	5.92	4-39	45.87
Minimum			8.43	6.12	0.45	1.72	28.43
Pepper shells:	1 4	1	1			1	_
Maximum			11.01	28.81	22.90	4.66	11.70
Minimum			7.00	7.82	0.79	1.53	9.28

	Ether :	Extract.		Nitr	ogen.	Total N
	Volatile.	Non-vola- tile.	Crude Fiber.	Total.	In Non- volatile Ether Extract.	non-vola- tile Ether Extract X 6.25.
Black pepper:						
Maximum	2.10	10.44	18.8q	2.38	0.45	13.12
Minimum	0.85	6.60	10.05	1.86	0.25	9.25
Average	1.30	7.67	11.12	2.11	0.31	11.20
White pepper:		1 ' '				
Maximum	1.66	7.26	7.65	2.14	0.34	11.56
Minimum	0.78	5.65	0.10	1.85	0.24	9.69
Average	1.17	6.46	4-17	1.97	0.30	10.44
Long pepper:	1					1
Maximum	1.01	7-53	10.01	2.04	0.22	12.06
Minimum	0.79	5.71	7.19	2.13	0.18	11.37
Pepper shells:		1		1	1	1
Maximum	1.11	4.67	28.22	1.82	0.12	11.25
Minimum	0.89	1.51	21.06	1.72	0.02	10.00
	!	1	1	I T	1	1

[†] Two samples of Acheen C pepper had a total ash of 8.00% and 8.04%, with 'ash insoluble in HCl' of 2.50% and 2.40% respectively. Eliminating these two samples, which were evidently abnormally high in sand and dirt, the highest total ash of the remaining 43 samples was 7.00%, while the highest ash insoluble in HCl was 1.80%.

Determination of Nitrogen in Black and White Pepper.—Winton, Ogden, and Mitchell have shown that the Kjeldahl and Gunning methods are inapplicable in the case of pepper, owing to the presence of piperin, but that the Gunning-Arnold † method gives accurate results. In accordance with this method, I gram of the sample is mixed with a gram each of copper sulphate and red oxide of mercury, about 16 grams of potassium

^{*} Mich. Dairy and Food Comm. Bul. 94.

[†] Zeits. anal. Chem., 31, 1892, p. 525.

sulphate, and 25 cc. of sulphuric acid in a Kjeldahl flask, for both digestion and distillation, of about 600-cc. capacity. The heating is conducted in the usual manner, beginning with a gentle heat till the frothing ceases, and gradually increasing the temperature till the mixture boils. The boiling is continued for three or four hours, after which the flask is cooled, and to it are added 300 cc. of water, 50 cc. of potassium sulphide solution,* and enough of a saturated solution of sodium hydroxide to render the reaction alkaline.

The flask is then connected to the condenser, and the distillation conducted as in the Gunning method (p. 69), using zinc dust to prevent bumping, receiving the distillate into standard acid, and titrating against standard alkali.

Nitrogen Determination in the Ether Extract.†—Ten grams of the sample are extracted with absolute ether for twenty hours in a continuous-extraction apparatus, the extract being collected in a tared Kjeldahl extraction- and distillation-flask, the same as used in the preceding section. The ether is then evaporated off, the residue dried to constant weight at 110° C. and its weight ascertained. The nitrogen is then determined in the ether extract by the Gunning-Arnold method.

Determination of Piperin.‡—Fifty grams of the sample are thoroughly exhausted with hot alcohol, and the alcohol extract evaporated to dryness. The dry residue is then treated with a solution of potassium hydroxide, and washed upon a filter. The residue is washed several times with the caustic alkali, which dissolves the resinous matters, and afterwards with water. It is then dissolved in alcohol, from which crystals of crude piperin separate on evaporation. These are redissolved in alcohol, and precipitated by the addition of water. The crystalline precipitate is collected on a tared filter, washed with water, dried, and weighed.

Piperin may be roughly estimated by multiplying the nitrogen in the ether extract by the factor 20.36.

The amount of piperin varies considerably, ranging in black pepper from 4 to 9 per cent.

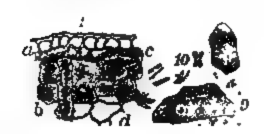
Microscopical Characteristics of Ground Pepper.—Moeller's representation of powdered black pepper shows what should be looked for under the microscope with the best conditions (Fig. 85). The shell of the peppercorn, a cross-section of which is shown at (1), consists of the

^{*} Forty grams K2S in I liter or water.

[†] Method of Winton, Ogden and Mitchell.

[‡] Villiers et Collin, Substances Alimentaires, p. 371.

epidermis, a, under which is a thin layer of brown parenchyma, c, while below this layer is shown the most characteristic portion of



the pepper shell, viz.: the thickened, colored, stone cells, b. These are as a rule inclined to be rectangular rather than rounded. At d is shown a bit of the colorless parenchyma of the fruit itself.

(2), (3), and (4) show a cross-section of the outer part of the berry, (2) representing the inner stone-cell layer, a single row of horseshoe-like cells, (3) the thin seed coat, and (4) the white perisperm, with its large cells. Here and there through the perisperm certain yellow contents are visible, consisting largely of resinous matter. A dark resin cell is shown at (4). The ethereal oil, starch, and piperin are found in this part of the berry.

Fig. 85.—Powdered Black Pepper under the Microscope. × 125. (After Moeller.)

(5) shows in plan view the mostly rectangular stone cells of the pepper shell, resting upon the epidermis (6). Groups of stone cells are frequently thus found with portions of the epidermis.

The inner rounded, or cup-shaped cells are shown in plan view at (7) and the seed skin at (8), masses of starch and separate starch granules are shown at (9), and crystals of piperin at (10).

The bast-parenchyma of the pepper stem is shown at (11), pieces of which are commonly found in powdered pepper, and (12) shows a fragment of one of the many-celled hairs which grow on the stem.

The rounded cup cells (7) are readily distinguished from the more rectangular stone cells (5). The walls of the cup cells are nearly always colorless, and the cells themselves empty.*

A water-mounted specimen of finely ground, black pepper, when viewed microscopically, shows most of the elements above described, at least in fragmentary form, though, in the case of the coarser particles,

^{*} The harder portions of the pepper, especially of the shell, are best examined by soaking for at least twenty-four hours in chloral hydrate, and mounting in this reagent on the slide.

by no means as clearly as by the use of chloral hydrate. Large polygonal masses of starch appear grouped as photographed in Fig. 256, Pl. XXXIV, if not rubbed out too fine under the cover-glass. Starch, indeed, is the most conspicuous element of pepper, being distributed more or less evenly throughout the mass. The powder may, however, be so finely reduced by abrasion under the cover-glass as to break up these starch masses wholly or in part, so that the granules may appear in much smaller groups or even singly. Fig. 255 shows such a field under a higher magnification. The individual granules of pepper starch average 0.003 mm. in diameter.

Besides the starch, and next to it the most numerous, one finds in the water-mounted black-pepper specimen many of the dark-yellow, thick-walled stone cells, patches of the colored parenchyma, and epidermis of the shell. Other elements of the perisperm, besides the starch, are seen in fragments, such as bits of resin, small droplets of oil, pieces of stems, and occasionally the needle-shaped crystals of piperin. Some of the rounded, cup-shaped cells are also usually found.

White pepper contains, of course, the same elements, but without the deeply colored stone cells and other characteristics of the shell, which has been removed from it.

Adulteration of Pepper.—The following U. S. standards for pepper have been adopted: For white pepper, non-volatile ether extract should not to be less than 6%; starch should not be less than 50% by the diastase method; total ash should not be more than 4%; ash insoluble in hydrochloric acid should not exceed 0.5%; crude fiber should not exceed 5%. One hundred parts of the non-volatile ether extract should contain not less than 4 parts of nitrogen. For black pepper, which should be free from added pepper shells, pepper dust, and other pepper by-products, non-volatile ether extract should not be less than 0%; starch by the diastase method should not be less than 0%; total ash should not exceed 0%; and crude fiber should not exceed 0%. One hundred parts of the non-volatile ether extract should contain not less than 0%. The adulterants used in ground pepper are many and varied.

Pepper Shells, which have been removed from the white pepper of commerce, are not infrequently ground and added to the cheaper grades of black pepper. When a sample of black pepper is shown by the microscope to contain more shells in proportion to the other elements than could be possible in a ground whole berry, added shells are indicated.

The analyst should, for comparison, grind in a mortar single berries of various grades, and familiarize himself with the appearance of the ground powder under the microscope, when the maximum amount of shells possible under natural conditions are present, noting especially the apparent number of stone cells of the outer coating. The familiar title of P. D. (pepper dust) originally given to ground pepper shells, stems, and "sweepings" is now applied in the trade not only to almost any cheap and appropriate material for admixture with pepper, but also, in a broader sense, to ground powder suitable as an adulterant for any spice.

The presence of pepper shells is indicated by an excess of ash, sand, and crude fiber, and a deficiency of starch.

Hilger and Bauer, also Hanus and Bien, advocate the determination of pentosans as a means of detecting pepper shells.

Ground Olive-stones constitute one of the most commonly found foreign materials used as an adulterant of pepper. The powder, sometimes called "poivrette," is very like white pepper in appearance, is wholly inert in taste, and thus forms an admirable adulterant. While best detected by their characteristic appearance under the microscope, the presence of ground olive stones may be shown by color tests with certain chemical reagents.

Pabst has adopted for this purpose a test first suggested by Wurster for the detection of wood pulp in paper. The reagent is prepared as follows: In a porcelain capsule 10 grams of commercial dimethyl anilin are mixed with 20 grams of pure concentrated hydrochloric acid, and at least 100 grams of cracked ice are added. Then, while stirring, a solution of 8 grams of nitrite of soda in 100 cc. of water are added little by little, and the mixture allowed to remain for half an hour, after which 30 or 40 cc. of hydrochloric acid are added, and 20 grams of tin-foil. The reduction is allowed to go on for half an hour, heating on the waterbath, if necessary. The tin is then precipitated by granulated zinc, the liquid is filtered, and the filtrate neutralized with carbonate of potassium or sodium to the point of forming a precipitate, the precipitate being dissolved by a few drops of acetic acid. Finally the volume is made up with water to 2 liters, adding, before doing so, 3 or 4 cc. of a concentrated solution of sodium bisulphite, to prevent oxidation. The reagent thus prepared will keep for several years in a brown, tightly stoppered bottle.

If a pinch of pepper, which contains ground olive stones, be heated gently with a little of the above reagent in a test-tube, the stone cells of the adulterant will be colored a bright red brown, and the colored particles will be seen to settle to the bottom of the tube, after shaking,

more quickly than the rest of the powder. Or, if the whole is poured from the test-tube into a porcelain dish, the color is more marked. Pure popper is not colored under this treatment with the reagent.

Jumeau uses for a color reagent 5 grams of iodine in 100 cc. of a mixture of equal parts of ether and alcohol. Enough of the finely ground pepper to be examined is placed in a porcelain capsule to cover the bottom of the dish, and sufficient iodine reagent is added to wet the entire mass, carefully avoiding excess. The thick paste is first mixed till homogeneous, and then allowed to dry in the air, after which it is broken up by a pestle, and the powder examined, either under the microscope, or by the naked eye. With pure pepper, a more or less deep-brown color is produced uniformly through the powder, but if olive stones are present, particles of these are colored yellow. With the naked eye as small an admixture as 2% of olive stones can thus be detected.

A solution of anilin acetate colors olive stones yellowish brown, while pure pepper appears grayish, or white.

Under the microscope olive stones are readily apparent, since the stone cells differ in size, form, and mode of grouping from those of pepper. Fig. 263, Pl. XXXVI, is a photograph of a water-mounted specimen of olive stones. They are for the most part entirely devoid of color, being long and narrow. In shape and manner of grouping they much resemble cocoanut shells (p. 419), but are distinguished from the latter from their lack of color.

Fig. 261 shows under low magnification a sample of pepper, bought on the market in Massachusetts, highly adulterated with olive stones. A large mass of the stone cells of the adulterant appears in the center of the field. Many of the stone cells are shown arranged end to end, so that what at first sight appear to be single, very long cells are in reality made up of several shorter ones. In ground olive stones one frequently finds, besides the stone cells, bits of the outer tegument of the seed, showing large cells with sinuous, rather thick walls; also bits of parenchyma, crossed frequently by fibro-vascular duct bundles.

Buckwheat Products.—Both the hulls and the middlings have been added to black pepper, and the middlings to white pepper. The starch of buckwheat possesses the added advantage, from the point of view of the spice-grinder, that it somewhat resembles pepper starch in microscopical appearance, not only in the shape of the starch granules, but also in the manner of grouping into masses. Compare Figs. 128 and 129, Plates II and III, showing buckwheat starch, with Figs. 255 and 256, Pl. XXXIV, respectively, showing pepper starch made under similar

conditions of magnification, etc. The starch granules and masses are coarser in the case of buckwheat than of pepper.

Fig. 260, Pl. XXXV, shows a photograph of a pepper sample adulterated with buckwheat, masses of both starches appearing in the same field.

Other Adulterants found in Massachusetts samples of pepper have been wheat and corn products, nutshells, cayenne, charcoal, turmeric, rice, sand, and sawdust. Charred cocoanut shells were at one time extensively used (see pp. 419 and 420).

Long Pepper, according to English analysts, has been used to a considerable extent as an adulterant. This is the fruit of the *Chavica Roxburghii*, a wild plant growing in India on the banks of rivers. The fruit, as its name implies, is long and cylindrical, while of about the same diameter as the spherical true peppercorns. Long pepper contains, as a rule, less than half the amount of piperin that true pepper does, and rather more starch than black pepper. Its taste is much less pungent than that of true pepper.

From its method of growth, long pepper is found with considerable dirt and sand adhering to the outer surface of the dried grains. This is due to the fact that the fruit often trails on the ground, and in gathering it the natives are not particular about removing the adhering soil. The surface of the fruit grains being very rough and irregular, much of the dirt remains dried thereon. The presence of long pepper thus materially increases the ash.

Long pepper possesses a very disagreeable, but peculiar odor, developed more especially when slightly warmed. For this reason, if for no other, it is not an ideal adulterant, since pepper containing it would not be palatable with warm food. At the present time it costs more than black pepper, and is used chiefly in mixed whole spices for pickles.

Brown gives the following analyses of samples of long pepper:

	Total Ash.	Sand and Ash Insol- uble in Hydrochlo- ric Acid.	Starch and Matters Converti- ble into Sugar.	Albumin- ous Matter Soluble in Alkali.	Cellulose.	Alcoholic Extract.	Ether Extract.	Total Nitrogen.
1	8.91	1.2	44.04	15.47	15.7	7-7	5-5	2.1
2	8.98	1.1	49-34	17.42	10.5	7.6	4-9	2.0
3	9.61	1.5	44.61	15.51	10.37	10.5	8.6	2.3

According to Brown and Heisch, the granules of long pepper starch under the microscope are larger than those of true pepper, and more angular. Stokes,* however, finds no such marked difference in the size

^{*} Analyst, XIII, p. 109.

of starch granules and his experience is shared by the writer. When the two specimens (long and true pepper) are viewed side by side in water mounts under the microscope, the average size of the long pepper-starch grains is a trifle larger than those of true pepper, though, unless compared directly, the difference is not readily apparent. Stokes suggests a method of distinguishing the two by polarized light. With crossed Nicols, so that a dark field is given, and with the specimen mounted in glycerin, true pepper starch shows an evenly dark appearance, using a low power, while with long pepper a "ghostly white" image is shown. Long pepper, when present in true pepper powder, may generally be rendered apparent by the development of the characteristic odor on heating. Bits of fluffy fiber from the catkin of the long pepper will always be found in the ground powder, and will be apparent under the magnifying-glass.

Microscopic examination of the crude fiber discloses the highly characteristic, large, beaded cells of the endocarp, also elements of the spindle.

RED PEPPER.

Nature and Composition.—According to the U. S. Standards red pepper is the red, dried, ripe fruit of any species of Capsicum, a genus of the nightshade family (Solanaceæ), indigenous to the American tropics, but now cultivated in nearly all warm and temperate countries, and is of two distinct kinds: cayenne pepper or cayenne, the dried ripe fruit of C. frutescens, C. baccatum, or some other small fruited species of Capsicum, and paprika, the dried ripe fruit of C. annuum, or some other large-fruited species of the genus, excluding seeds and stems.

Cayenne is characterized by its extreme pungency and the small size of the pods, which seldom exceed 2 cm. in length. The leading commercial varieties are Zanzibar and Japan, the latter being the more brilliant in color.

"Capsicums" or "Bombay Chillies" are low grade peppers of a brown color, with pods 2 to 3 cm. long, which now are said to come from the vicinity of the river Niger in Africa.

Paprkia is a variety of C. annuum grown in Hungary. The powder is of a deep red color and has a sweetish, mildly pungent flavor.

Pimiento is a large-fruited pepper grown in Spain. The succulent pericarp is much used for stuffing olives while the dried pod is ground as a spice, often being substituted for the more valuable Hungarian varieties. The kitchen garden peppers, of which over thirty varieties are cultivated in the United States, also belong to the species C. annuum.

The capsicum plant has solitary flowers, with a five-cleft corolla, and the fruit is of an elongated, conical form. The surface of the fresh fruit is smooth and very red, but it loses some of its brilliance in drying, and becomes shriveled. The pericarp is thin and tough, and at its base is a five-lobed calyx, greenish brown in color, terminating in a thick stem. The fruit proper is divided into two or three cells, which are separate and distinct at the lower portion, but which unite and form one at the top. The cells inclose a large number of yellow, wrinkled, kidney-shaped seeds, containing a fleshy endosperm, and a curved embryo.

Red pepper contains a fixed, bland oil, found in both pod and seed, but more abundantly in the latter, considerable resinous and mucilaginous material, a red coloring matter confined to the pod, and the active principle capsicin, a crystalline alkaloid, to which much of the pungency is due. The capsicin is present in both seeds and pod, but is more abundant in the latter, where it is dissolved in the oil.

Capsicin may be isolated, according to Thresh, by extracting powdered cayenne with petroleum ether, mixing the red residue left on evaporating off the solvent with two or three times its weight of oil of almonds, and exhausting the mixture with alcohol. On evaporating the alcohol extract, the capsicin crystallizes out in narrow, thin plates, very soluble in alcohol, but insoluble in water. They volatilize at 100°, and condense in small drops.

The red coloring matter is soluble in ether, petroleum ether, carbon bisulphide, and chloroform, but sparingly soluble in alcohol.

Analyses of Cayenne.—Richardson * gives the following data of analyses of two pure samples of cayenne:

	Water.	Ash.	Fixed Oil.	Volatile Camphor, etc.	Fiber.	Albumin- oids.	Undeter- mined.	Total.	Nitrogen.
A B	2-35 5-74	9.06 5-24	0.12	26.99 17.90	16.88 18.10	13.13	41.47 40.24	100	2.10

Maximum and minimum data of ash and non-volatile ether extract of fourteen samples of cayenne, sold in sealed packages in Connecticut, and analyzed by Winton and Mitchell are as follows:†

	Ash.	Non-volatile Ether Extract.
Maximum	7.18 5.88	19.14 15.59

^{*} U. S. Dept. of Agric., Div. of Chem., Bul. 13, p. 211.

[†] An. Rep. Conn. Exp. Sta., 1898, p. 175.

Winton, Ogden, and Mitchell * analyzed eight samples of whole chillies, representing three varieties, namely Zanzibar, Japan, and Bombay, the summarized results being as follows:

			Ash.		Ether I	Extract.
	Moisture.	Total	Soluble in Water.	Insoluble in HCl.	Volatile.	Non-vola- tile.
Maximum	7.08 3.67 5-73	5-96 5-08 5-43	4.93 3.30 3.98	0.23 0.05 0.15	2-57 0-73 1-35	21.81 17.17 20.15
	Alcohol Extract.	Reducing Matters as Starch, Acid Con- version.	Starch by Diastase Method.	Crude Fiber.	Nitrogen, ×6.25.	Total Nitrogen.
Maximum	27.61 21.52 24.35	9.31 7.15 8.47	1.46 0.80 1.01	24.91 20.35 22.35	14.63 13.31 13.67	2.34 2.13 2.18

The percentages of "starch by the diastase method" given in the above table represent errors of the process as neither cayenne or paprika contain an appreciable amount of starch.

Analyses of Paprika and Pimiento.—Doolittle and Ogden † have made exhaustive analyses of known samples of Hungarian and Spanish red pepper, including determinations of non-volatile ether extract, and iodine number of this extract, which are of especial value in detecting added oil. A summary of their results is given on page 442.

Microscopical Structure of Red Pepper.—Fig. 86, from Moeller, shows the appearance under the microscope of various elements of powdered paprika. (1) is a sectional view through the outer portion of the fruit shell or pod, showing the epidermis a, and beneath this the collenchyma layer. The inner epidermis is shown at (2), with its cells thick-walled in places, and inclosing brilliant, red oil drops of coloring matter. (3) represents the outer, and (4) and (5) the inner epidermis in plan view. The outer epidermis of cayenne, which is the element of chief value in distinguishing this from paprika, is shown at (6).

A cross-section through the seed shell is shown at (7), a being the epidermis of the seed, b the parenchyma layer directly beneath, and c the tissues of the endosperm. (8) shows in plan view the peculiar seed

^{*} Am. Rep. Conn. Exp. Sta., 1898, pp. 200-201.

[†] Jour. Am. Chem. Soc., 30, 1908, p. 1481.

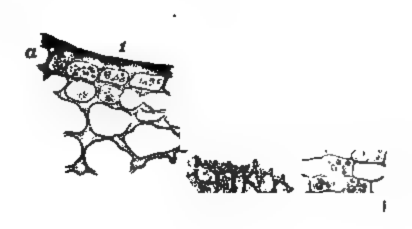
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	mples.			Ash.	1	Alkalinity o	nity of h.*	Ether Extract.	xtract.	Non-	ters by raion.		٠٤٤٠
	Number of Sa	O °oo1 ta seoJ	Total.	Soluble in Water.	Insoluble in HCl.	.latoT	Water- soluble.	Volatile.	Non-volatile.	lodine No. of volatile Eth Extract.	Reducing Mat Acid Conver as Starch.	Crude Fiber.	o× ,nagoniN
Whole Pods:													
Hungarian: Maximum	•	9.39	90.4	2.68	0.23	8.8	2.60	1.25	11.99			19.83	17.44
Minimum		7.26	5 63	4.67	0.05	6 22	4.07	0.17	7.42	130-3	19.50	15.10	14.06
		8.54	6.28	5.12	0.10	7.05	4.04	0.85	9.30		20.44		15.43
Spanish: Maximum	a	8.58	62.9	5.79	90.0	7.00	5.70	1.18	10.39			15.37	16.87
Minimum		8.28	5.24	4.59	0.05	6.10	4.10	1.18	10.39	134-5		15-37	14.62
Average		8-43	6.02	5.19	0.05	7.00	8	1.18	10.39			15.37	15.75
Pods (Seeds, placentæ and stems removed):				-									
Hungarian: Maximum.	7	10.86	6.9	6.10	0.08	8.60	6.10	1.10	6.69	138.0	24.52	23.61	15.37
Minimum		9.45	5.50	4.85	0.03	8.90	4.85	0.44	4.01			16.66	
Average		10.37	6.03	5-44	0.05	6.92	5-53	0.80	5.08	133.1	23.90	19.50	16.33
Spanish: Maximum	т	10.35	2.68	6.68	0.30	8.8 8	و. 8	1.40	4.76		19.96	15-19	
Minimum		6.26	6.23	5.70	0.05	7.65	5.70	0.51	4.48			14.80	11.64
Average		69.4	7.17	6.22	0.11	8.45	6.35	0.95	4.62	131.0		15.00	13.05
Seeds and Placentæ:													
Hungarian: Maximum	7	6.46	4.93	3.72	6.00	6.80	3.80	8.	22.34	135.4	18.16	20.11	21.19
Minimum		5.8	3.06	1.72	0.05	3.40	1.50	0.95	17.66			17.29	
Average		5.80	3.8	2.85	0.07	4.85	2.71	1.50	20.31			18.74	
Spanish: Maximum	3	6.58	5.30	4-35	0.11	4.40	3.00	2.25	19.80			24.01	16.25
Minimum.		5.19	3.41	2.23	0.04	3.90	1.80	1.56	18.99		16.12	19.48	15.50
Average		5-74	4.41	3-37	90.0	4.17	2.33	16.1	19.40	132.1	16.12	21.74	15.92
Stems:													
Hungarian, Average	7	6.55	11.32	8.36	0.44	14.07	8.46	0.48	1.94	15.88	:	24.47	15.87
Spanish, Average	8	4-73	15.50	13-09	0.26	14.10	0.11	0.29	%	:	:	29.99	11.56
			- 		1								

• Cc. N HCl per gram of material, using methyl orange as indicator.

epidermis, the appearance of which Moeller compares with that of intestines. At (9) is shown one of the isolated cells of this epidermis more highly magnified, while (10) shows the epidermis of the calyx.

Figs. 211 and 212, Pl. XXIII, show photomicrographs of powdered cayenne. In Fig. 211 is shown a large bit of the outer epidermis of the fruit pod, while in Fig. 212 appears a smaller portion of this same kind of epidermis, and next to this the characteristic skin of the seed shell, with its striking markings suggestive of the convolutions of the intestines. Yellow or yellowish-red droplets of oily coloring matter are distributed through the field. Starch grains are absent.



 \boldsymbol{a}

b

Fig. 86.—Powdered Cayenne under the Microscope. X125. (After Moeller.)

Adulteration of Red Pepper.—The U. S. standards for cayenne are the following: Non-volatile ether extract should be not less than 15%; total ash should not exceed 6.5%; ash insoluble in hydrochloric acid should not exceed 0.5%; starch by the diastase method should not exceed 1.5%, and crude fiber should not exceed 28%.

The most common adulterants of cayenne are the starches of the cereal grains, corn and wheat. Ground pilot bread and crackers are especially common. Besides these the writer has found in the routine examination of cayenne samples in Massachusetts, ginger, nutshells, turmeric, rice, gypsum, buckwheat, olive stones, mustard hulls, ground redwood, red ocher, and coal-tar dyes. Fig. 213, Pl. XXIV, shows a sample adulterated with wheat, corn, and cocoanut shells.

Mineral Adulterants, such as gypsum, and red other and other pigments, are all to be looked for in the ash by methods of qualitative analysis. An abnormally high ash is suggestive of adulteration. According to Vedrodi, the ash of genuine cayenne should not exceed 5.96. The presence of red other is rendered apparent by the high content of iron.

Salts of lead and mercury are rarely if ever now used for color.

Ground Redwood.—Numerous varieties of redwood are commonly used to intensify the color of cayenne, especially when otherwise highly adulterated with colorless materials, such as the starches. The redwood is sometimes used alone, and sometimes in mixture with turmeric. Both redwood and turmeric are readily recognized under the microscope.

Fig. 214, Pl. XXIV, shows a cayenne sample adulterated with corn starch and red sandalwood, a mass of the latter filling the center of the field. The wood fibers of the dyestuff, even when finely ground, are very striking under the microscope, showing a brick-red color.

Detection of Coal-tar and Vegetable Colors.—Oil-soluble coal-tar and vegetable colors may be tested for in cayenne and paprika by an adaptation of Martin's butter-color method, shaking the ether extract of the sample with the alcohol and carbon bisulphide mixture, page 535. The carbon bisulphide dissolves the oil and natural color, while the overlying alcohol layer holds in solution many of the artificial coloring matters that may be employed.

The natural colors of cayenne and paprika are sparingly soluble in alcohol, but readily soluble in carbon bisulphide. The separated alcohol is examined for colors by methods given elsewhere.

Tests for coal-tar dyes should also be made by Sostegni and Carpentieri's, or Arata's method (p. 793).

Szigeti * treats the suspected sample with water acidified with acetic acid, and boils in this solution a bit of wool, which, if carotin or a coal-tar dye be present, is colored red. If the color is carotin, it will be removed

^{*} Zeits. landw. Versuchs. Oesterreich, 5, 1902, pp. 1208, 1222.

from the wool by treatment with petroleum ether, or by heating at 100° C. for some hours, but if a coal-tar dye, it will still remain fixed thereon.

Detection of Added Oil in Paprika.—The color of paprika is often intensified by grinding with olive oil or some other oil. This form of adulteration is detected by determinations of non-volatile ether extract and iodine number of the extract.

Doolittle and Ogden's results (page 442) indicate that paprika prepared from the pods after removal of the stems, seeds, and placentæ should not contain more than 7.0% of non-volatile ether extract and the iodine number of this extract should not be less than 127, while that made from the whole pod should not contain over 12.00% of non-volatile ether extract, and the iodine number should not be less than 130.

The method employed by Doolittle and Ogden consists in extracting 2 grams of the material on a dried filter paper with anhydrous ether, collecting the washings in a tared, glass-stoppered flask, distilling to remove ether, and drying the residue for thirty minutes, or to constant weight, in a steam bath. Iodine number is determined in this non-volatile ether extract by the Hanus method (page 491).

This method gives a lower percentage of non-volatile ether extract, and a higher iodine number than extraction in a continuous apparatus for 16 hours. The latter method is open to the objection that, owing to the presence of slowly soluble resins, the results are appreciably affected by the rate of extraction.

The following method gives concordant results, although somewhat different from those by the preceeding methods: Dry 4 grams of the material in a watch glass over sulphuric acid for 18 hours, place in a 100 cc. flask, add anhydrous ether to the mark, and shake at five minute intervals for 30 minutes. Filter through a dry paper into a flask, keeping the funnel covered with a watch glass to prevent evaporation. Pipette off an aliquot of the filtrate equivalent to about 0.2 gram of extract, distil off the ether, dry, and determine iodine number as above described.

GINGER.

Nature and Composition.—Ginger as a spice is the ground rootstock of the Zingiber officinale, an annual herb of the family Zingiberaceæ, growing to a height of from 3 to 4 feet. It is a native of India and China, but is cultivated quite extensively in tropical America, Africa, and Australia.

The root is dug when the plant is a year old, and when the stem has

withered. If the root, when freshly dug and scalded to prevent sprouting, is dried at once, it forms the so-called black ginger, of which Calcutta and African are the common varieties. When decorticated, the product is known in commerce as white ginger, the chief varieties being Jamaica, Cochin, and Japan. The best variety is Jamaica ginger. The scraped root is sometimes bleached to make it still whiter, or sprinkled with carbonate of lime.

In commerce whole or black ginger appears in "hands" 4 to 10 cm. long, and from 10 to 15 mm. in diameter. These usually have three or four various-sized, irregular branches, some short and thick, others elongated. The epidermis is gray or yellowish gray in color, more or less wrinkled, and beneath it is a reddish-brown layer. The inner portion of the dried root is white or yellowish. The root is hard, and of a compact, horny structure.

White or decorticated ginger appears in "hands" of smaller diameter than the black, and yields a lighter colored powder on grinding. Preserved ginger root is prepared by boiling the root in water, and curing with sugar or honey. Much of the preserved ginger comes from Canton.

The distinguishing features of ginger are its large content of starch, its volatile oil, and its resinous matter. Inasmuch as the epidermis contains a large amount of pungent resin, it is easy to see how the peeled or decorticated variety is inferior.

Oil of ginger is very aromatic, and of a greenish-yellow color. Its specific gravity ranges from 0.875 to 0.885. It is slightly soluble in alcohol. Of its composition little is known.

Richardson's analyses in full of five samples of whole ginger-root are as follows:

Water.	Ash.	Volatile Oil.	Fixed Oil and Resin.	Starch.	Crude Fiber.	Albumin- oids.	Undeter- mined.	Nitrogen.
9.60 9.41 10.49 11.00 10.11	3-39 3-44	1.84 2.03	4.07 2.29 3.04	49-34 53-33 50.58 49-34 50.67	2.05 4.74 1.70	7.00 10.85 9.28	13.44 18.91 15.58 19.21 11.66	1.12 1.74 1.48

Summaries of Winton, Ogden, and Mitchell's analyses of eighteen samples of whole ginger, representing the common white and black varieties, as well as of two samples of exhausted ginger, are as follows:

			Ash.			Ether E	xtract.
	Moisture.	Total.	Soluble in Water.	Insoluble in HCL	Lime (CaO).	Volatile.	Non-vola- tile.
Ginger: Maximum Minimum Average. Lxhausted ginger from English ginge ale works Exhausted ginger from extract works	10.6	3.61 4 5.27 1 2.12	4.09 1.73 2.71 0.59 3.55	2.29 0.02 0.44 0.18 1.50	3-53 o.20 o.80	3.09 0.96 1.97 1.61 0.13	5.42 2.82 4.10 3.86 0.54
	Alcohol Extract.	Keducing Matters by Acid Con- version, as Starch.	Starch by Diastase. Method.	Crude Fiber.	Nitrogen, ×6.25.	Cold-water Extract.	Total Nitrogen.
Ginger: Maximum Minimum Average Exhausted ginger from English ginger-ale works. Exhausted ginger from extract works.	6.58 3.63 5.18 4.88 1.52	62.42 53-43 57-45 59.86	60.31 49.05 54.53 54.57	2.37 3.91	9-75 4-81 7-74 6-94	17-55 10.92 13.42 6.15 16.42	0.77

McGill * records the analyses of ninety-eight samples of ground ginger as sold in the Canadian market. Of thirty-two of these, pronounced pure on analyses, the following is a summary:

				• Ash.					
	Moisture or Loss on Dry- ing at 100°.	Petro- leum- ether Extract.	Cold- water Extract.	Total.	Soluble.	Insoluble.	Alkalin- ity of Soluble Ash as K ₂ O.		
Maximum	12.00 9.50	6.13	15.48	7.84 3.67	3.15	3-99	.133		

According to Vogl, the proportion of ginger ash varies quite widely according to the kind, but should never exceed 8%.

Exhausted Ginger and Methods of Detection.—There are two kinds of exhausted ginger commercially available for admixture with ground spice, as an adulterant. One is the product left after extraction with strong alcohol in the making of extract of Jamaica ginger, and the other the residue from extraction with either very dilute alcohol, or with water,

^{*} Dept. Inl. Rev. Canada Bul. 48, pp. 10, 11.

in the manufacture of ginger ale. Ground, exhausted ginger is rarely substituted wholly for the pure variety, since, from its lack of pungency, the sophistication would be too apparent. It is rather used to mix with the latter in varying proportions, and as an adulterant of other spices.

Ginger that has been exhausted by extraction with alcohol has been deprived of most of its volatile oil, which is found in the "extract," while for the manufacture of ginger ale, a water extract, or at most a very dilute alcoholic extract is best adapted. Such a water extract does, as a matter of fact, remove much of the valued pungency, so that the residue, or exhausted ginger, is rather inert.

Either the alcohol- or the water-extracted variety of exhausted ginger, when present in considerable amount, would be apparent, one by the alcohol and ether extract, and the other by the abnormally low coldwater extract, and water-soluble ash.

Dyer and Gilbard * first called attention to the water-soluble ash as a reliable means of indicating exhausted ginger. Six samples of ginger of known purity were analyzed by them, their results being summarized as follows:

		Total Ash.	Water- soluble Ash.	Alcohol Extract, after Ether Extract.
Pure ginger (6 samples):	Highest	4.1	3-	3.8
	Lowest	3.1 3.8	1.9	2.1
Exhausted ginger (6 samples):	Highest	2.3	0.5	1.5
	Lowest.	1.1	0.2	
	Average	1.8	0.35	1.2

Allen and Moor † pointed out the value of the cold-water extract as a help in detecting exhausted ginger, especially when taken in connection with the soluble ash, showing that the presence of this adulterant is assured, when the soluble ash is as low as 1%, and the cold-water extract is less than 8%.

Determination of Cold-water Extract.—Winton, Ogden, and Mitchell's Method.‡—Four grams of the ground sample are placed in a 200-cc. graduated flask, and the latter is filled to the mark with water, and shaken at half-hour intervals during eight hours, after which it is allowed to

^{*} Analyst, XVIII (1893), p. 197.

[†] Analyst, XIX (1894), p. 194.

[‡] U. S. Dept. of Agric., Bur. of Chem., Bul. 65, p. 59; Bul. 107 (rev.), p. 164.

stand at rest for sixteen hours in addition. The contents are then filtered, and 50 cc. of the filtrate evaporated to dryness in a platinum dish. It is then dried at 100° to constant weight and weighed.

Microscopical Structure of Ground Ginger.—Fig. 87, from Moeller, shows elements of ginger root, from which the epidermis has not been



Fig. 87.—Powdered Ginger under the Microscope. X125. (After Moeller.)

removed. A bit of the large-celled cork (or dead protective tissue of the epidermis) is shown in plan view at (1); at (2) is shown in cross-section the parenchyma in which the starch is contained, h being an oil-cell; (3) shows the parenchyma in longitudinal section, with bast fibers. Fragments of spiral ducts are shown at (4), and starch grains at (5). (6) is a cross-section in the extreme interior of the root.

The most prominent feature of powdered ginger is the starch grains (5), which Moeller compares in shape to tied sacks.

Fig. 228, Pl. XXVII, is a photomicrograph of pure, ground ginger, mounted in water, showing the starch grains inclosed in the cells of the parenchyma. Fig. 231 shows the starch grains alone. The granules of ginger starch are ellipsoidal, and as a rule very clear and transparent, being for the most part entirely devoid of either hilum or concentric rings.

Occasionally granules are to be found, however, with faint concentric markings, and even with an apparent hilum. The characteristic form of the ginger starch granule is more or less egg-shaped, with a small protuberance near one end. This protuberance serves to readily distinguish the starch granules of ginger from those of wheat, with which ginger is frequently adulterated. While wheat granules are of various sizes, the grains of ginger starch are as a rule much more uniform.

Adulteration of Ginger.—U. S. standard ginger should meet the following requirements: Starch by the diastase method should not be less than 42%; crude fiber should not exceed 8%; total ash should not exceed 8%! lime should not exceed 1%; ash insoluble in hydrochloric acid should not exceed 3%.

Besides exhausted ginger, the most common adulterants found in powdered ginger are turmeric, wheat, corn, rice, and sawdust. Sawdust of soft wood is a not uncommon adulterant, and care should be taken to distinguish between the wood fiber natural to the ginger root, and that of the foreign variety. A careful study should be made of finely ground, soft-wood sawdust, with its long spindle cells and lateral pores, as shown in Fig. 266, Pl. XXXVII, and the wood fiber of the genuine ginger root. A large admixture of sawdust would materially increase the percentage of crude fiber.

Fig. 234, Pl. XXIX, shows a sample of ginger adulterated with corn and wheat. Fig. 232 shows a mass of wheat bran in an adulterated sample.

Fig. 233 shows ginger adulterated with turmeric.*

TURMERIC.

Nature and Composition.—Turmeric, while largely used as an adulterant of other spices (especially of ginger and mustard), possesses some value as a condiment in itself, forming, for instance, the chief ingredient of curry powder.† Turmeric (Curcuma longa) belongs to the same family (Zingiberaceae) as ginger, having a perennial rootstock, and an annual stem. It is a native of the East Indies and Cochin-China. Its chief ingredients are starch, a volatile oil, a yellow coloring matter (curcumin), cellulose, and gum.

^{*} This photomicrograph is very disappointing, in that it fails to show the intense yellow of the central mass of turmeric.

[†] Curry powder consists of a mixture of turmeric, cayenne, and various pungent spices.

Curcumin (C₁₄H₁₄O₄) is insoluble in cold water, but readily soluble in alcohol. It is extracted from powdered turmeric by boiling the latter with water, filtering, and extracting the residue with boiling alcohol. The alcoholic solution is filtered, evaporated, and the residue extracted with ether. The ether extract contains the curcumin, together with a small amount of volatile oil.

Curcuma oil is an orange-yellow, slightly fluorescent liquid, its specific gravity being 0.942.

The following analyses of turmeric were made in the writer's laboratory:

Variety.	Mois- ture.	Total Ash.	Ash Soluble in Water.	Ash Insoluble in HCl.	Total Nitrogen.	Protein, N×6.25.	Total Ether Extract.
China	9.03 9.08 8.07	6.72 8.52	5.20 6.14 4.74	0.11	1.73 0.97 1.56	10.81 6.06	10.86 12.01 10.66
Average	8.73	5•99 7•07	5.36		1.42	9-75 8.88	11.17

Variety.	Volatile Ether Extract.	Non-vol- atile Ether Extract.	Alcohol Extract.	Crude Piber.	Reducing Matter by Acid Con- version, as Starch.	Starch by Diastase Method.
China.	2.01	8.84	9.22	4.45	48.69	40.05
Pubna	4-42	7.60	7.28	5.84	50.08	29.56
Alleppi	3.16	7.51	4-37	5.83	50.44	33.03
Average	3.19	7.98	6.96	5-37	49-73	34.21

Microscopical Structure of Turmeric.— Moeller's representation of characteristics of powdered turmeric is reproduced in Fig. 88. The epidermis is shown at (1) with one of the numerous, one-celled hairs that grow from it, also the scar left after one of the hairs has been removed; (2) shows in plan view the cork immediately under the epidermis. The tender-celled parenchyma is shown in cross-section at (3), and in longitudinal section at (4). In some of the cells of the parenchyma are found dark-yellow lumps of resin (h), and vascular ducts (g), but by far the most numerous and striking contents of the parenchyma-cells are the bright-yellow masses of "paste balls" (3a) and the starch granules, one of which is shown in (3). See also Plate XIII. The starch grains in the water-mounted powder show under the microscope in masses, usually of a deep-yellow color, unless very finely rubbed out, when they appear for the most part in fragments.

The whole starch granule appears somewhat in the form of a clamshell, with very distinct markings. When fragments of the starch granules are carefully examined, these distinct markings are so strongly characteristic, even in the smallest pieces commonly found in the powdered sample, as to nearly always serve to identify them. See Fig. 171, Pl. XIII.

Turmeric as an Adulterant.—Turmeric is a material especially adapted by its deep-yellow color to intensify mustard and ginger, especially when

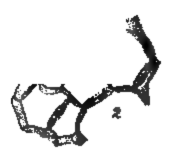


Fig. 88.—Powdered Turmeric under the Microscope. X125. (After Moeller.)

these spices are adulterated with the lighter-colored cereal starches, hence it is very commonly found in these spices, both with and without other adulterants.

It is also frequently used in small quantities in adulterated cayenne, mace, and various spices, to counteract the colors of other dyestuffs, such as ground redwood, which in itself would sometimes be too intense if used alone.

Turmeric, when present to any marked extent in a powdered spice, may be detected chemically, by extracting the material with alcohol, pouring off the latter, and soaking in it a piece of filter-paper. Turmeric, if present, will stain the latter yellow, turning red with alkali, especially apparent after drying. Soak the yellow paper in a solution of borax, acidulated slightly with hydrochloric acid. When dry, a rose-red color will indicate turmeric, turning dark olive when dilute alkali is applied.

MUSTARD.

Nature and Composition.—Mustard is the seed of the mustard plant, an annual belonging to the family *Cruciferæ*, and to the genus *Sinapis*, or *Brassica*, as it is sometimes called. The plant is an herb, native throughout Europe, and cultivated extensively in the United States. It grows to a height of from 3 to 6 feet, having yellow flowers and lyrate leaves.

Two varieties commonly used are Brassica or (Sinapis) alba, white mustard, and Brassica (or Sinapis) nigra, black mustard, the ground spice being as a rule a mixture of the two. In the trade these varieties are known as brown and yellow mustard respectively. The seeds of both varieties are globular, those of the black mustard being small, and of a dark-brown color on the outside and yellow within. White mustard seeds are considerably larger than the black, being pale yellow in color on the outside.

The surface of the black mustard seeds is reticular, and full of small depressions, while the white variety is much smoother. There are several layers forming the husk of the seed of both varieties, and within the husk is the yellowish-colored kernel or embryo, with two cotyledons.

Both black and white mustard contain from 31 to 37% of fixed oil, a soluble ferment known as myrosin, and a sulphocyanate of sinapin. Mustard seeds contain no starch, and very little volatile oil as such. Black mustard seed contains sinigrin, or myronate of potash (not found in the white seed), which, when moistened with water, forms by hydrolysis the volatile oil of black mustard, otherwise known as allyl isothiocyanate, in accordance with the following equation:

$$\begin{array}{c} KC_{10}H_{10}NS_2O_9 + H_2O = C_0H_{12}O_0 + C_2H_5CNS + KHSO_4. \\ \begin{array}{cccc} Potassium & Mustard & Potassium \\ myronate & oil & bisulphate \end{array}$$

Mustard Oil (volatile) is a colorless, or slightly yellow, highly refractive liquid of a very strong odor, and capable of blistering the skin when

brought in contact with it. It is optically inactive. Its specific gravity varies between 1.016 and 1.030. It boils between 148° and 156°. It turns reddish brown by exposure to light.

Volatile oil of black mustard forms thiosinamine with ammonia, as follows:

$$C_3H_5CNS + NH_2 = CS.NH_2.NH.C_3H_5.$$

Thiosinamine is soluble in hot water, from which it crystallizes in tufts of monoclinic crystals, having a melting-point of 74° C. It is precipitated by silver nitrate, mercuric chloride, and Mayer's solution.

White mustard differs from the black in containing a sulphur compound, sinalbin, C₃₀H₄₂N₂S₂O₁₆. This is a glucoside. Sinalbin by hydrolysis forms an oil of white mustard, in a somewhat similar manner to the potassium myronate of black mustard, and according to the following equation:

$$\begin{array}{ccc} C_{30}H_{42}N_2S_2O_{15} + H_2O = C_7H_7ONCS + C_6H_{12}O_6 + C_{16}H_{24}NO_5HSO_{4^*}\\ & \text{Sinalbin} & \text{Sinalbin} & \text{Glucose} & \text{Sinapin acid sulphate} \end{array}$$

Sinalbin Mustard Oil cannot be obtained by the distillation of white mustard, being sparingly volatile with steam.

Sinalbin mustard oil somewhat resembles that from black mustard, being quite as pungent, but less strong in odor when cold. It is soluble in dilute alkali.

Fixed oil of mustard is a bland, tasteless, and nearly odorless oil, its specific gravity at 15° varying between the limits of 0.914 to 0.918. It is said to be used to some extent as an adulterant of table oils, being separated by pressure from the crushed mustard seeds before the latter are ground into "flour." The chief use of mustard oil is in mixture with other oils as an illuminant.

MUSTARD FLOUR.—In the process of preparing the ground spice commonly known as mustard "flour," the seeds are first crushed and separated by winnowing from the hulls, the latter being incapable of the fine grinding necessary to produce a smooth flour. The yellow hulls are, however, found in the cheaper grades of ground mustard, and both varieties of hull are frequently used in the wet mustard preparations, sold in bottled form. In order to produce an even, dry powder, free from lumps, it is necessary to remove a large portion of the fixed oil, which is indeed of no value in the final product, and this is done by subjecting the crushed material to hydraulic pressure, during which process the

mustard is molded together into thin, hard plates, called "mustard cake." This is then broken up and reduced to fine powder by pounding. Richardson's * analyses of whole-seed flour, prepared by himself without the removal of the fixed oil, are as follows:

	Water.	Ash.	Volatile Oil.	Fixed Oil.	Starch.	Crude Fiber.	Albumin- oids.	Undeter- mined.	Nitrogen.
White seed	5.57 3.33 6.17 4.83 4.11 3.11 4.62	4.29 5.23 4.99 5.96 4.88 4.07 5.61	-97 1.84 -55 1.27 1.35 2.06	33.56 34.83 28.12 31.96 36.63 31.51 39.55	.00	5.40 9.05 9.50 8.50 16.18 6.90 10.84	28.88 25.56 23.44 31.13 24.69 30.25 25.88	27.23 16.35 12.16 22.10	4.09 3.75 4.98 3.95 4.84

Winton and Mitchell made no full analyses of mustard seed of known purity, but the following is a summary of analyses of 18 samples of commercial mustards, sold in packages in Connecticut, and not found to be adulterated:

	Total	Ether 1	Extract.	Reducing Matters by Acid	Starch by	Crude	Nitrogen	
	Ash.	Volatile.	Non-vol- atile.	Conver-	Diastase Method.	Fiber.	×6.25.	
Maximum	7-35 4.81 5-99	1.90 0.00 0.56	28.10 17.14 20.61	6.12 1.85 4-33	2.08 0.28 1.07	4.87 1.58 2.58	43.56 35.63 39.57	

The following analyses of 5 samples of mustard flour, 6 samples of mustard hulls, and 6 samples of whole mustard, were made in the author's laboratory in 1903:

^{*} U. S. Dept. of Agric., Div. of Chem., Bul. 13, part 2.

Reduc'g Mi ters by Di stase.	22000	.37			200 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	-
Reduc's Ma ters by Ac Conversion	20 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	9.13	S. S	1 0 m	0 0 4 4 0 1 0 4 0 4 4 0 1 0 4	13.82
Crude Piber	4 26 3 15 3 30 3 57 2 57 2 50 2 57	3 30			# # # # # # # # # # # # # # # # # # #	80 0.0
Total Ash.	**************************************	6.75	W40-40.	t to us	2000000 200000 200000 200000 200000	6.83
Total Nitra Sen	# 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	00 6		•	65 77 75 88 66 57 75 88 66 57 88 67 75 88 68 88 88	7.09
Alcohol Es	32.88 20.64 20.71 30.71	39 S.C.	PP-1-00	F-4 (f)	######################################	24.83
Reducing by Diestar	**************************************	90	a nomente	94 6	172	1.48
Reducing by Acid Calon.	# 4 2 4 6 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8	6.85	0	-0.0	24420400 44420400	8.62
Crude Fiber	80 K 8 K K K 8 8 8 8 8 8 8 8 8 8 8 8 8 8	2.43	Bun 4400		44440 4 N.O. 44	\$ 04
Volatile Oil	8 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4	:			2.000 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	:
Total Nitro	00 -0 -00 00 -0 -00 00 - 0 -00 00 - 0 -00 00 -00 -	6.78	1 1	+	44484844	4: 4 I
Alcohol Ex	2 - 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	24 30	4-404-06	, n, o	13 98 13 98 14 75 14 98 16 91 16 91 16 91	15.50
Non-volatile Extract.	110 2 2 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3	18.50			2 0 0 0 2 4 4 0 X	31.22
Volatile Btl tract.	0000000	0.0			0000000	
Total Bihe	17 40 10 20 10 28 11 2 65 12 65 19 65	r8.59	,		33-8-30-30-30-30-30-30-30-30-30-30-30-30-30-	31.22
Agh Insolu	90 H N M H W W	. 27	********	0 H	************	6
Water-solul	420254	61.	0,4 - 40 - 40	, o . c	044.0 24.0 24.0 24.0 24.0 24.0 24.0 24.0	.56
Total Ash.	พุลุพพุลลล	. e.s		* *	40044444 0000464000	4 9.5
Moisture.	22.024.00	6.96	******	4 W	2000 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	6.37
	"flour": commercia lish brown forma brown man brown v. of brown	waneties emoved flour: 1	hulls	Av. of yellow hulls verage of all samples of hulls ground mustare	if brown fiornia brown to of brown needs tran yellow glish yellow flornia yellow if ornia yellow if ornia yellow	
	Total Ash. Total Ash. Total Lincolul Anter-solul Trotal Lincolul Trotal Lincolul Alcohol Est Crude Fiber Sen Total Nitro Crude Fiber Crude Fiber Total Nitro Crude Fiber Sen Sen Total Nitro Crude Fiber Conversion Conversion Conversion Conversion Conversion Conversion Conversion	Meter-solution of the first of	Condensity of the Prince Ball Mitton brown flours. Total Mitton bro	Moisture, Mois	Moisture. Mois	Av. of brown flours.

* Freed from hulls and with a portion of the fixed oil removed.

† These hulls necessarily have a little of the inner seed adhering thereto.

‡ Ground to a fine paste with full content of oil and hulls.

Piesse	and	Stansell	give	the	following	composition of	mustard ash:
		~	8				

,	White	Seeds.	Brown Seeds.
	Yorkshire.	Cambridge.	Cambridge.
Potash. Soda. Lime Magnesia Iron oxide Sulphuric acid Chlorine	21.29 0.18 13.46 8.17 1.18 7.06	18.88 0.21 9.34 10.49 1.03 7.16 0.12	21.41 0.35 13.57 10.04 1.06 5.56 0.15
Phosphoric acid. Silica. Sand Charcoal.		35.00 1.12 1.95 15.14	37.20 1.41 1.38 7.57

Determination of Myronate of Potassium and Sinapin Sulphocyanate.*

—Extract at least 50 grams of the powdered material with several portions of a mixture of equal parts of water and alcohol, digesting with the aid of heat in a flask with a return-flow condenser. Evaporate the alcoholic extract in a tared dish to dryness, and heat at 105° to constant weight. After weighing, incinerate the residue at a temperature sufficiently high to transform to the neutral sulphate the potassium bisulphate resulting from the decomposition of the myronate. The weight of myronate of potassium is obtained by multiplying the weight of neutral sulphate (the final ash) by the factor 4.77. This, deducted from the total weight of the dried alcoholic residue as above, gives that of the sulphocyanate of sinapin.

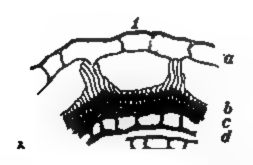
Determination of Mustard Oil in Mustard Flour.—Roeser's Method.†
—Mix 5 grams of the sample with 60 cc. of water and 15 cc. of 60% alcohol, and let stand for two hours. Distil into a flask containing 10 cc. of ammonia, and, after about two-thirds of the solution have been distilled off, mix the ammoniacal distillate with 10 cc. of tenth-normal silver nitrate solution, and allow the mixture to stand for twenty-four hours, after which make up with water to 100 cc. Filter, and treat 50 cc. of the filtrate with 5 cc. of tenth-normal potassium cyanide solution. Titrate the excess of cyanide with the tenth-normal silver nitrate, using as an indicator a 5% solution of potassium iodide, made slightly ammoniacal.

^{*} Girard et Duprée Analyse des Matières Alimentaires, p. 678.

[†] Abs. Analyst, XXVII, 1902, p. 197.

The percentage of mustard oil present is found by multiplying by 2 the number of cubic centimeters of silver nitrate solution taken up by the oil, and multiplying this product by the factor 0.3137.

Microscopical Characteristics of Powdered Mustard.—The principal features of powdered black mustard are represented in Fig. 89. The



seed shell or hull is shown in cross-section at (1), a being the polygonal-celled epidermis, b a layer of palisade-shaped cells, and c a thin pigment layer, the brown coloring matter of which is colored blue by iron salts; d is the alcurone layer and obscure parenchyma, and e the small-celled tissue of the cotyledons, containing fixed oil and albumen.

- (2) shows in plan view the various layers of the seed shell, the letters of reference corresponding to those of (1).
- (3) shows in plan view a bit of the extreme outer mucilaginous layer of the seed-hull.

Fig. 247, PI XXXII, shows the ap-

mustard. This is a photomicrograph of the ground hulled seed without the extraction of the oil, and should not be taken as a standard for commercial mustard "flour," from which, as a rule, a large portion of the oil has been removed. The cellular tissue of the mustard shows in the form of granular masses of loose, fine gray texture; the globular bodies are oil drops. Here and there through the field of ordinary ground mustard are to be seen patches of the yellowish layer of the seed skin of the brown mustard, a mass of which is shown in Fig. 248, with dark-brown spots distributed regularly through it. This is the layer shown at (2) b, Fig. 89. The hull of the yellow seed, also common in powdered mustard, is similar in appearance, having dark-

Patches of the outer hull layer represented by (3) in Fig. 89 are also very common in the commercial mustard flour. Mustard contains no starch.

brown spots, but with nearly colorless or gray cell walls, instead of yellow.



Fig. 89. — Powdered Mustard under the Microscope. ×125. (After Moeller.)

Adulteration of Mustard.—U. S. standards for mustard are as follows: Starch, by diastase method, should not exceed 2.5% and total ash should not exceed 8%.

It is difficult to draw the line between the amount of mustard hulls which may naturally occur in ground mustard, and the excess amount which is sometimes added as an adulterant. Samples in which the patches of hulls predominate in number over the regular cellular tissue of the seed, as seen under the microscope, are undoubtedly adulterated by the fraudulent admixture of ground hulls, that have been separated out from the crushed mustard seeds intended for higher grades. Samples of mustard flour thus adulterated are common.*

In determining starch in mustard, it should be borne in mind that mustard hulls have considerable reducing matter by the diastase process.

The most common adulterants of mustard, other than excess of hulls, are wheat, turmeric, millet and other weed seed, and rice. Yellow, oil-soluble azo-dyes are also employed.

Other adulterants' found in Massachusetts have been potato starch, cayenne, corn, and gypsum or "terra alba" (the latter being found in one instance to the extent of 21%).

Fig. 250, Pl. XXXIII, shows a sample of mustard adulterated with wheat bran. Very little besides the adulterant appears in this field.

The common practice of adulterating mustard with wheat is an outgrowth of the old notion that a certain amount of wheat flour was necessary to prevent lumping.

Samples of cheaper mustard flour are occasionally found to contain small amounts of wheat and foreign starch, apparently of accidental occurrence. This is undoubtedly due to the fact that in some localities wild mustard often grows in the wheat-fields, so that after the wheat crop has been harvested, the mustard is gathered and sold. Such mustard seed would naturally contain varying admixtures of wheat, and sometimes seeds of various weeds, which it would not be profitable to separate out, even

^{*}It is claimed by some manufacturers that the hulls thus removed are not used as an adulterant of cheaper mustard flours, in view of the fact that it is difficult or impossible to grind them finely enough, but that they are used up in the manufacture of compound mustard pastes. A sample of ground mustard was recently found by the writer, in which it was noticed that a large number of yellow lumps were distributed through it. These lumps were picked out, transferred to the microscope slide, and crushed and rubbed out under the cover-glass. Examined under the microscope, they were found to consist entirely of a mixture of mustard hulls and turmeric, which would seem to show that hulls were present in this case as an adulterant.

if it were possible to do so. Such contaminated mustard enters into the manufacture of the cheapest grades of flour only.

Fig. 249, Pl. XXXIII, shows the flour of the Dakota brown mustard, or charlock (*Brassica Sinapistrum*), which is the commonest of the wild mustards. This species is characterized by the presence in the pallisade cells of a dark brown substance, which on treatment under the cover glass with chloral hydrate solution assumes a beautiful blood-red color. This highly characteristic reaction is hastened by gentle heating, and permits the ready detection of charlock in mustard flour.

Detection of Coloring Matter.*—Turmeric is best detected by the microscope (see pp. 451 and 452). Oil-soluble coal-tar dyes should be tested for as in the case of cayenne. Nitro colors, such as naphthol yellow (Martius yellow) and naphthol yellow S, are detected by dyeing tests, with subsequent examination of dyed fabric according to the scheme on p. 799.

PREPARED MUSTARD.—This product consists of a mixture of ground mustard seed or mustard flour with salt, spices, and vinegar. The U. S. standards require that it should contain not more than 24% of carbohydrates calculated as starch, not more than 12% of crude fiber, and not less than 35% of protein.

Most of the product consumed in the United States is of domestic manufacture, although until the passage of the federal food law it was customary to designate it German or French mustard, or label it in a foreign language.

Composition and Adulteration.—The common admixtures are wheat flour, maize flour, and other starchy matter, mustard hulls, sugar, chemical preservatives, and artificial colors.

Of 28 brands examined in Connecticut in 1905 by Winton and Andrew,† 13 contained cereal flour (wheat or corn), 4 salicylic acid, and 25 artificial color (turmeric, nitro-color or azo-color). A summary of the

^{*}Recently some very yellow samples of powdered mustard have appeared on the market that are apparently free from foreign color. Their method of manufacture is kept secret. From the fact that they contain nearly, if not quite, the full content of fixed mustard oil that would be present if the oil had not been previously expressed, and for various other reasons, it is probable that the color is due largely to the presence of the fixed oil, which has a deep-yellow color, and which has hitherto been generally removed for purposes of fine pounding and to avoid caking.

In such samples, the oil, previously pressed out, is, after pounding, restored, and with it much of the color. Incidentally in such a process oil-soluble coal-tar dyes may conveniently be dissolved in the mustard oil, in order to intensify the color, and the analyst should be on the outlook for such foreign colors.

[†] An. Rep. Conn. Exp. Sta., 1905, p. 123.

analyses of those brands free from cereal flour and those containing it follows:

		In the Material as Sold.									
	Water.	Acidity Calculated as Acetic Acid.	Total Solids.	Total Ash.	Com- mon Salt.	Ash other than Salt			Reduc- ing Matters by Acid Conver- sion, as Starch.	Ex- tract.	Pat.
Prepared mustard free from cereal flour: Maximum Minimum Average	73.01		23.67 13.32 18.36	2.60	1.78		3.62	0.77	2.92 1.83 2.40	4.21	7.23 2.12 4.12
Prepared mustard con- taining cereal flour: Maximum Minimum	85.63 70.44	3 · 54 1 · 86	27.70 9.89	4.21	3·39 1.51	1.16 0.48	6.38 1.53	1.59	13.69	15.35 3.82	3 · 25 0 · 76

	In the Dry, Pat, and Salt-free Material.					
	Ash.	Protein.	Crude Fiber.	Reducing Matters by Acid Conver- sion, as Starch.	Nitrogen- free Extract.	
Prepared mustard free from cereal flour: Maximum Minimum Average	10.66 7.35 8.94	43.94 32.01 39.44	14.12 7.77 9.89	24.37 16.82 20.11	44.76 34.98 41.73	
Prepared mustard containing cereal flour: Maximum Minimum	9.68 4.84	33.89 21.37	18.44 0.45	59.22 24.51	66.42 41.79	
Whole mustard seed (analysis by the author. See page 456): Maximum Minimum Average	7.64 6.28 6.83	48.31 37.50 44.31	10.33 7.24 8.05	15.91 11.94 13.82	48.55 37.84 40.81	

The following methods for the analysis of prepared mustard were used by Winton and Andrew, and afterwards adopted by the Association of Official Agricultural Chemists:

Solids, Ash and Salt are determined in one portion of 5 grams of the thoroughly mixed material, following the usual methods. The salt is calculated from the percentage of chlorine.

Ether Extract.—Ten grams of the material and about 30 grams of sand are placed in a capsule, and dried on a water bath with stirring. The dried residue is ground and extracted with anhydrous ether in the usual manner.

Reducing Matters by Acid Conversion are determined directly in the material, without previously washing, by the method described on page 411.

Crude Fiber.—Eight grams of the material (equivalent to about 2 grams of dry matter) are treated in the usual manner (page 277), except that care is taken to add at first only a small amount of the 1.25% acid or alkali, and shake thoroughly until all lumps are broken up, as otherwise these lumps will resist the action of the solution and the results will be high.

Protein.—Nitrogen is determined by the Kjeldahl or Gunning method, and the result multiplied by 6.25.

Dyes and Preservatives are detected by the methods described in chapters XVII and XVIII.

NUTMEG AND MACE.

Nature and Composition.—Both nutmeg and mace occur in the fruit of several varieties of trees of the genus Myristica, especially of the Myristica jragrans or Myristica moschata, belonging to the family Myristicaceæ. The nutmeg tree is a native of the Malay archipelago, and grows from 20 to 30 feet high, somewhat resembling the orange tree in appearance. It does not produce flowers till its eighth or ninth year, after which it bears fruit constantly for many years. The fruit is a globular, pendant drupe, about 5 cm. in diameter, of a yellowish-green color, the pericarp of which, when ripe, splits in two, showing within it the kernel, completely surrounded by a fleshy, fibrous aril, or covering of a crimson color. This covering, when dried, furnishes the mace of commerce, while the inner kernel, which is a hard, brown seed, is the nutmeg.

The nutmeg seed or kernel, when gathered, is surrounded by a thick tegument, marked with depressions corresponding to the lobes of the aril or mace, and by a second thin, inner envelope, closely adhering to the seed. The whole seed is dried in the sun for about two months, or by the aid of heat, the tegument becoming separated from the kernel, and, by breaking with a hammer, is readily removed. The kernels are then commonly washed in milk of lime, and again dried, or they are sometimes treated with dry, powdered, air-slaked lime. Liming is alleged to prevent sprouting and ward off the attacks of insects. The so-called brown nutmegs of commerce are those which have not been treated or coated with lime.

MUTMEGS are spheroidal, sometimes nearly spherical, from 20 to 25 mm. long and 15 to 18 mm. in diameter. The outer surface is somewhat furrowed. A cross-section of the kernel shows the grayish-brown, starchy endosperm, mottled with the dark-brown, resinous veins of the perisperm. These veins on pressure with the finger nail present an oily appearance. Near the end of the nutmeg which is attached to the stem, is a small cavity, in which is the undeveloped embryo with two cotyledons.

Nutmeg contains a considerable amount of fixed oil, a volatile oil, starch, and albuminous matter. Its volatile oil is colorless, and is soluble in three parts of strong alcohol. The specific gravity of nutmeg oil varies between 0.865 and 0.920, and its specific rotary power $(a)_{D}=14$ to 28.

Richardson's analyses of three samples of nutmeg are as follows:

	Water.	Ash.	Volatile Oil.	Pixed Oil or Fat.	Starch, etc.	Crude Fiber.	Albu- minoids.	Nitro- gen.
Whole limedGround limed		3-27 2-22 3-15	2.84 3.97 2.90	34-37 37-30 30.98	36.98 40.12 41.77	11.30 6.78 9-55	5.16 5.42 5.25	.83 .87 .84

König gives the following minimum and maximum composition of nutmeg:

l	Minimum.	Maximum.
Water.	4.2	12.2
Albuminoids	15.2	6.1
Volatile oil	2.5	4.0
Fat	31.0	37-3
Carbohydrates	29.9 6.8	37·3 41.8
Cellulose.	6.8	12.0
Ash	2.2	3-3

Winton, Ogden, and Mitchell analyzed four samples of nutmeg of known purity, the following being maximum and minimum results:

			Ash.		Ether 1	Extract.
	Moisture.	Total.	Soluble in Water.	Insoluble in HCl.	Volatile.	Non-vola- tile.
Maximum		3.26 2.13	1.46	0.01	6.94 2.56	36.94 28.73

	Alcohol Extract.	Reducing Matters by Acid Con- version, as Starch.	Starch by Disstase.	Fiber.	Nitrogen × 6.25.	Total Nitrogen.
Maximum	17.38	25.60 17.19	24.20 14.62	3-72 2-38	7.00 6.56	1.12

Microscopical Structure of Nutmeg. (Fig. 90.)—The thin-walled cells of the parenchyma of the endosperm or albumen are shown at



Fig. 90.—Powdered Nutmeg under the Microscope. ×125. (After Moeller.)

(1), with starch grains. Simple and compound granules of the starch are shown at (2). Aleurone grains appear as shown at (3), and (4) represents in plan view the epidermis, or brown seed coat, with its numerous layers of flat cells. Powdered nutmeg under the microscope in watermount shows most commonly a spongelike, loose meshwork of bruised or broken cellular tissue, with many starch granules, and occasional fragments of the epidermis.

Fig. 240, Pl. XXX, is a photomicrograph of a water-mounted sample of pure nutmeg. The starch granules of nutmeg are different from other starches in appearance, being almost circular as a rule, quite uniform in size (averaging 0.006 mm. in diameter), and having very distinct central hyla.

Adulteration of Nutmeg. — The U. S. standards for nutmegs are as follows: Non-volatile ether extract should be not less than 25%; total ash should not exceed 5%; ash insoluble in hydrochloric acid should not exceed 0.5%; crude fiber should not exceed 10%.

This spice is more often sold in the whole form, since the house-wife much prefers to grate the whole nutmeg, rather than to use the ground material. It is hence less liable to adulteration than the other spices, though of late more of the ground nutmeg is being sold in packages. Samples of ground nutmeg have been found in Massachusetts adulterated with wheat and nutshells. One sample was found to contain at least 25% of ground cocoanut shells.

Nutmegs which have become mouldy, or have been eaten out by insects, have been imported for grinding, as sound nutmegs are not readily reduced to a powder. Such a product is obviously unfit for consumption.

An inferior variety is known as the Macassar nutmeg. This lacks much of the agreeable pungency of the better grades.

MACE.—The crimson-colored aril that surrounds the nutmeg kernel within the pericarp, as above described (p. 462), has many narrow, flattened lobes. In the process of drying to form the mace of commerce, it loses its brilliant red color, and turns a yellowish brown. When dried, it is brittle, somewhat translucent, and of a pungent odor. Whole mace appear on the market in the form of flat membranous masses, 3 to 4 cm. long.

It contains no starch as such, but has a modified form of starch known as amylodextrin. This is a carbohydrate, $C_{30}H_{62}O_{31}+H_2O$, which produces with iodine a red coloration. Mace has a large amount of fixed oil, as well as considerable resinous and albuminous matter, and a volatile oil which much resembles that of nutmeg.

The specific gravity of volatile oil of mace is rather higher than that of nutmeg oil. Its specific rotary power, $(a)_n = 10$ to 20.

König's figures for the composition of mace are as follows:

	Minimum.	Maximum.
Water	4.9	17.6
Albuminoids	4.6	6.1
Volatile oil	4.0	8.7
Fat	18.6	29.1
Carbohydrates	41.2	44.I
Cellulose	4.5	8.9
Ash	1.6	4.1
Alcoholic extract	45.1	55-7

Richardson gives the following as the results of analyses of three samples made by him:

	Water.	Ash.	Volatile Oil.	Resin.	Unde- ter- mined.	Crude Fiber.	Albu- minoids.	Nitro-
Whole mace	4.86	4.10 2.65 2.20	4.04 8.66 8.68	27.50 29.08 23-33	41.17 35.50 34.68	8.93 4.48 6.88	4-55 6.13 5.08	·73 .98 .81

Winton, Ogden, and Mitchell's analyses of four samples of pure Banda or Penang mace, as well as of Bombay and Macassar mace, are summarized as follows:

1	Maintana	Ach.				Ether Extract.		
	Moisture.	Total.	Soluble in Water.	Insoluble in HCl	Volatile.	Non-vols- tile.		
True mace: Maximum	12.04 9.78 11.05 4.18 0.32	2-54 1.81 2.01 2.01 1.98	1.33 1.06 1.13 1.11	0.21 0.00 0.07 0.03 0.07	8.65 6.27 7.58 5.89 4.65	23.72 21.63 22.48 53.54 59.81		
	Alcohol Extract.	Reducing Matters by Acid Con- version, as Starch.	Starch by Diastase.*	Crude Piber	Nitrogen ×6.25.	Total Nitrogen		
True mace: Maximum Minimum Average Macassar Bombay (adulterant)	24.76 22.07 23.11 32.89 44.27	34.42 26.77 31.73 10.39 16.20	30.43 23.12 27.87 8.78 14.51	3.85 2-94 3.20 4-57 3.21	7.00 6.25 6.47 7.00 5.06	I.12 I.00 I.03 I.12 0.81		

^{*} The figures in this column do not express starch, but amylo-dextrin, which like starch may be determined by the diastase method.

Microscopical Structure of Mace.—Fig. or shows characteristics of mace, (1) being a cross-section through it, (2) a plan view of the

epidermis, showing its elongated, often nearly rectangular cells, and (3) the largecelled parenchyma, in which are numerous oil globules. The contents of the parenchyma cells are for the most part colorless, consisting of protein, fat, granules of amylodextrin, which are shown at (4). At (5) are shown fragments of vascular tissue.

the Microscope. X125. (After Moeller.)

The water-mounted powder of pure mace shows no highly colored fragments, Fro. 91.-Powdered Mace under but as a mass, is white or grayish, and of loose texture. Occasional pale, yellowish, lumpy masses appear, and pale-

brown fragments of the seed coating. The amylodextrin granules (which are colored red-brown by solution of iodine) are very small.

Adulteration of Mace.-U. S. standard mace should contain not less than 20 nor more than 30% of non-volatile ether extract; nor more than 3% of total ash; nor more than 0.5% ash insoluble in hydrochloric acid; nor more than 10% of crude fiber.

Turmeric and cereal starches are not uncommonly found in mace, but by far the most common adulterant is the so-called false, or wild mace, otherwise known as Bombay mace.

Bombay Mace (Myristica jatua) is almost entirely devoid of odor or taste, being nearly as inert as so much starch. It is most properly regarded as an adulterant from its lack of pungency, even though in a sense it is a variety of mace.

Its non-volatile ether extract is twice as high as that of Penang mace, and at room temperature the fixed oil of Bombay mace is a thick and viscous fat, while that of Penang and other maces is a thin oil.

The refractive indices of the fixed oils of various species of pure, as well as of Bombay mace, as determined by Lythgoe in the writer's laboratory, are as follows:

	7	D at 35° C.
Banda Mace	(1)	1.4848
"	(2)	1.4747
"	(3)	1.4829
Batavia Mace	(1)	1.4893
"	(2)	1.4975
Papua Mace	(1)	1.4816
"	(2)	1.4795
West Indian Mace	(1)	1.4766
Bombay Mace	(1)	1.4615
"	(2)	1.4633

The microscope indicates at once when Bombay mace is present in a sample. The oil glands situated in the outer layers of Bombay mace are very strongly colored, and contain a deep-red, resinous substance, very different from anything to be found in true mace. The glands of the more interior layers of wild mace have, moreover, a balsam-like substance of a bright-yellow color. In powdered Bombay mace, when mounted in water, nearly every field shows both the red and the yellow lumps in considerable number.

Hejelmann's Test for Bombay Mace * consists in saturating a strip of filter-paper with an alcoholic solution of the mace, and removing the excess of liquid by pressure between filter-paper. On treating with a

^{*} Pharm. Zeit., 1891.

drop of dilute sodium or potassium hydroxide solution, a red color is produced in presence of the wild mace.

Waage's Test.—One part of the mace is extracted with ten parts of alcohol, and potassium chromate solution is added to the extract. If Bombay mace is present, the solution becomes red, and the precipitate, which is at first yellow, becomes red on standing. True mace gives a yellow solution and precipitate, and the latter does not change greatly on standing.

Turmeric is tested for chemically as on p. 789.

Macassar Mace is sometimes designated as wild mace, but it is by no means as inert as the Bombay variety, and possesses a wintergreen-like odor. Its taste, while distinctive, is not that of true Penang mace. It is distinctly an inferior article, and its volatile oil content, as shown by the analyses on p. 466, is considerably below the minimum for true mace.

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(See also References on the Microscope, p. 98.)

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CHAPTER XIII.

EDIBLE OILS AND FATS.

Nature and Properties.—The oils and fats are the glycerin salts or glycerides of the fatty acids, the most important, on account of their occurrence in nearly all fats and oils, being the triglycerides of oleic, palmitic, and stearic acids, known as olein, palmitin, and stearin, respectively.

Fats and oils are insoluble in water, and are almost insoluble in cold 95% alcohol, though they are somewhat soluble in absolute alcohol. They are readily soluble in ether, petroleum ether, chloroform, amyl alcohol, oil of turpentine, and carbon bisulphide.

Following is a list of the fatty acids whose glycerides are found in edible oils and fats, together with their melting- and boiling-points when these have been determined, and the oils and fats in which they occur.

ACIDS OF THE ACETIC SERIES CaHanO.*

Name.	Formula.	Mdting- point.	Boiling- point.	Occurrence in Oils and Pats.
Butyric† Caproic† Caproic† Capric† Lauric. Myristic Palmitic Stearic. Arachidic Behenic Lignoceric	C ₃₀ H ₄₀ O ₃ C ₃₀ H ₄₀ O ₃ C ₃₁ H ₄₀ O ₃	-6.5° 16.5 31.3 43.6 53.8 62.6 69.3 77 83-84 81	162.3 200 236 268-270 176 196.5 215 232.5	Butter, cocoa butter. Butter, cocoanut oil. Cocoanut oil, cocoa butter. sesame oil. Nearly all oils and fats. Fats and nearly all oils, excerolive and corn. Peanut, olive (trace), rape (trace Rape, mustard. Peanut.

^{*} Lewkowitsch, Oils, Pats, and Waxes, 3d ed. (1904), pp. 63-71.

[†] These four soids are the only once that can be distilled under ordinary pressure without becoming decomposed.

ACIDS	OF	THE	OLEIC	SERIES	CaH	O~
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Name.	Formula.	Melting- point.	Boiling- point.	Occurrence in Oils and Pats.
Hypogæic Oleic Iso-oleic* Rapic		33° 14° 44-45° 33-34°	236° 232-5° 264°	Peanut. Nearly all fats and oils. Rape and mustard.

ACIDS OF THE LINOLEIC SERIES CaH2a-4O2

Name.	Formula.	Melting- point.	Boiling- point.	Occurrence in Oils and Pats.		
Linoleic	C ₁₈ H ₃₂ O ₂	Under—18°C.		Olive, cottonseed, peanut, sesame, cocoa butter, poppy seed, sun-flower.		

^{*} Solid oleic acid.

Saponification of Fats and Oils.—By this term is meant the decomposition of the glycerides composing the fats or oils, whereby the triatomic alcohol glycerin and the fatty acids are separated. The saponification process is commonly applied in carrying out many determinations of value on fats and oils, such as those of the soluble and insoluble fatty acids, the Reichert value, etc. As commonly carried out, the triglycerides are first split up into glycerin and the soluble soaps of the fatty acids by the action of caustic alkali, usually in solution in alcohol. This part of the process in the case of a given oil, composed, for example, of stearin, olein, and palmitin, is illustrated as follows:

(1)
$$C_3H_5(C_{18}H_{35}O_2)_3 + 3KOH = C_3H_5(OH)_3 + 3K(C_{18}H_{35}O_2)$$

Stearin or triglyoeryl Potassium stearate stearate

(2)
$$C_3H_5(C_{16}H_{31}O_2)_3 + 3KOH = C_3H_5(OH)_3 + 3K(C_{16}H_{31}O_2)$$
Palmitin or triglyceryl polmities

(3)
$$C_3H_5(C_{18}H_{23}O_2) + 3KOH = C_3H_5(OH)_3 + 3K(C_{18}H_{33}O_2)$$
Olein or triglyceryl oleate
Oleate

Potassium oleate

These "soaps," or potassium salts of the fatty acids, are further decomposed by the action of sulphuric acid into the free fatty acids and potassium sulphate, in the case of potassium stearate, as follows:

$$2K(C_{18}H_{35}O_2) + H_2SO_4 = K_2SO_4 + 2H(C_{18}H_{35}O_2)$$

Potassium stearate

ANALYSIS OF EDIBLE OILS AND FATS.

No class of food products presents more difficulties to the analyst than the fats and oils, in that the various physical and chemical constants by which one derives information as to their nature or purity differ within such wide limits that it is not easy to prescribe absolute standards. Many elements enter in to cause this variation, chief among which are, in vegetable oils, the large number of varieties of fruits or seeds from which each oil is in different localities obtained, as well as the various grades of each oil with respect to refining. In the animal fats, butter and lard, the kind of food fed to the animal undoubtedly influences the constants of the fat, and in all fats and oils much depends upon their age, and the conditions under which they are kept as to temperature, exposure to moisture, light and air, etc.

Rancidity should not be confounded with acidity, although rancid oils usually are high in acids. Lewkowitsch holds that fatty acids are liberated by the action of moisture in the presence of enzymes. If in addition the oil is exposed to air and light, the fatty acids are acted on causing rancidity, which is detected by taste and smell, although chemically little understood. As a rule rancidity develops more readily in liquid oils in which olein predominates than in solid fats. To avoid changes samples should be kept in a dark, cool place in tight containers.

Judgment as to Purity of a given oil or fat should not be hastily given. It is sometimes comparatively easy to prove the presence and approximate amount of an adulterant, the various constants all serving to identify it without fail. Again, in some cases it is easy to pronounce the sample adulterated, without being able to definitely state the exact nature of the adulterant. The tests to be employed depend on the particular case in hand. Sometimes the determination of two or three constants will be sufficiently definite.

Again, a large number of tests must be made before one can intelligently form an opinion. It should be borne in mind that skilful manufacturers may adulterate the edible oils and fats with mixtures intended to confuse the chemist, and yield on analysis constants that are entirely misleading.

Much information may usually be gained by carefully noting the color, taste, odor, and appearance of the sample.

Filtering, Measuring, and Weighing of Fats.—These manipulations naturally present some difficulties in the case of solid fats not encountered with liquid oils.

A steam- or hot-water-jacketed funnel as represented in Fig. 92 is convenient for filtering fats, or, in the absence of this contrivance for keeping the fat in a molten condition, a hot funnel may be employed, the filtering being best conducted in a warm closet or oven.

Portions of the fat for the various determinations may be measured off with a pipette while the fat is still hot, but a much better way is to

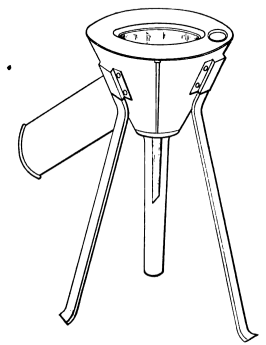


Fig. 92.-Jacketed Funnel for Hot Filtration.

cool the fat (over ice if necessary), and to weigh the desired amounts in the solid state. This can very readily be done by placing a flat platinum or other dish on the scale-pan, covering it with a moderately thick, cut filter-paper somewhat larger in diameter than the dish and designed to lie flat upon it, and taking the tare of both dish and filter. The solidified fat, after mixing with a stirring-rod, is transferred in one or more portions to the middle of the filter, and the exact weighed amount is obtained, after which, by carefully handling the edges of the filter and folding in the latter, the fat with the filter may be transferred to a flask or other receptacle.

Specific Gravity.—The specific gravity of liquid oils is most conveniently taken either at room temperature or at 15.5°, being always

best referred to the latter. Either the hydrometer, Westphal balance, Sprengel tube, or pycnometer are employed, according to the degree of accuracy required. If taken at any other temperature than 15.5° , say at room temperature, T, the specific gravity may be computed at 15.5° by the formula

$$G = G' + K(T - 15.5),*$$

in which G is the specific gravity at 15.5°, G' the specific gravity at T° , and K a factor varying with the different oils as follows:

Oil,	Correction for 1° C.	Observer.		
Cod-liver oil. Lard oil. Olive oil. Peanut oil. Rape oil. Sesame oil. Cottonseed oil.	.000658 .000629 .000655 .000620 .000624	A. H. Allen C. M. Wetherill C. M. Stillwell A. H. Allen		

FACTORS FOR CALCULATING SPECIFIC GRAVITY.

Unless the most accurate work is necessary, it is sufficient to assume in all cases K = 0.00064, in which case the formula becomes G = G' + 0.00064(T - 15.5).

In the case of solid fats, it is most convenient to take the specific gravity of the melted fat. This may be done at any temperature above the melting-point by either of the instruments above described, or at the temperature of boiling water by the Westphal balance or pycnometer.

When the pycnometer is used, it is immersed in a water-bath, the temperature of which is well above the melting-point of the fat, say 35° or 40°. While still immersed nearly to the neck, it is carefully filled with the melted fat and kept in the bath till the fat has acquired the same temperature, usually about fifteen minutes. If the pycnometer is provided with a thermometer stopper, this will serve to indicate the temperature; otherwise a separate thermometer is inserted in the bath. The pycnometer is then removed, cleaned, dried, and cooled to the room temperature, at which it is weighed. The factors employed in the above formula for calculation of specific gravity of solid fats at 15.5° are as follows:

^{*} Allen, Com. Org. Anal., 3d ed., vol. 2, pt. 1, p. 33.

FACTORS	FOR	CALCULATING	SPECIFIC	CDAVITY

Fats.	Correction for 1° C.
Cocoa butter Tallow. Lard. Butter fat Cocoanut stearin. Cocoanut oil. Palm nut oil.	0.000717 .000675 .000650 .000617 .000674 .000642

Either the Westphal balance or the hydrometer may be used directly on the melted fat, carefully recording the temperature and calculating as above.

For making the determination with the Westphal balance at the temperature of boiling water, the melted fat is contained in a vessel immersed in a boiling water-bath, and kept sufficiently long to acquire that temperature, which is carefully noted.

A. O. A. C. Method.*—The pycnometer, being perfectly clean, is first weighed with the stopper, after which it is filled with freshly boiled, hot, distilled water and placed in a bath of boiling water, where it is kept for half an hour, replacing any loss by evaporation in the flask with boiling distilled water. The stopper of the pycnometer, previously heated at 100°, is then inserted, and the flask removed and wiped perfectly dry. It is then allowed to cool nearly to room temperature, and weighed on the balance when the temperature is the same as that of the room.

The flask, being again perfectly clean and dry, is filled while hot with freshly melted and filtered hot fat, free from air-bubbles, and kept for half an hour in a boiling water-bath, after which the stopper, previously heated as before to 100°, is inserted, and the flask taken from the bath and wiped dry. It is then allowed to cool and weighed when the temperature of the room has been reached.

The specific gravity is calculated by dividing the weight of the fat by the weight of the water previously found.

Having once obtained the weight of the flask and the weight of a volume of water contained therein when at boiling temperature, these figures can be constantly used without redetermination, if the flask is cleaned thoroughly each time.

Calculation of Proportions of Two Known Oils in Mixture.†—This may be roughly accomplished from the specific gravity of the mixture and of the oils known to compose it.

^{*} U. S. Dept. of Agric., Bur. of Chem., Bul. 65, p. 21.

[†] Villiers et Collin, Les Substances Alimentaires, p. 646.

Le. G = specific gravity of mixture, D and D' = specific gravity of the two oils, and X = % oil of specific gravity D. Then $X = \frac{100(G - D')}{D - D'}$.

EDIBLE OILS AND FATS ARRANGED IN ORDER OF SPECIFIC GRAVITY

	Specific Gravity.
Cocoa butter	.976 to .950
Mutton tallow	-953 '' -937
Beef "	.952 '' .943
Butter	.940 '' .926
Lard	.938 '' .934
Poppyseed oil	.927 " .924
Sunflower oil	.926 '' .924
Corn oil.	.926 '' .921
Cottonseed oil	.925 " .922
Sesame oil.	.924 " .923
Peanut oil.	.921 " .917
Mustard oil	.920 " .916
Olive oil.	.918 " .916
Rape oil	.917 " .913

The Viscosity, or degree of fluidity in the case of edible oils, is of less importance than in the case of lubricating oils, and gives little insight into the nature or purity of the sample.

Hence a discussion of various viscosimeters and their use will not be included here, but reference is made to Lewkowitsch* for information regarding them.

The Refractive Index, and the reading on the arbitrary scale of the butyro-refractometer, express in two different and interchangeable terms the refraction value, a useful constant of fats and oils and easily determined.

For the routine examination of fats and oils the butyro-refractometer is more convenient than the Abbé refractometer, and the readings obtained by the former instrument are less cumbersome than refractive indices.

These instruments and details with regard to their manipulation are described in Chapter VI.

The readings on the scale of the butyro-refractometer may be readily transformed into refractive indexes and vice versa by table or by means of the Leach and Lythgoe slide rule (page 107). Lythgoe's† table on pp. 478 and 479 is useful as showing readings on the butyro-refractometer of all the edible oils and fats at various temperatures.

^{*} Chemical Analysis of Oils and Fats, 3d ed., 1904, pp. 197-209.

[†] Tech. Quarterly, 16, 1903, p. 222.

CALCULATED READINGS ON BUTYRO-REFRACTOMETER OF EDIBLE OILS AND FATS.

Temp. C.			Cacao Butter.	Beef Lard Tallow. Stearin.		Beef Oleo.	Lard.†	Lard Oil.	
45.0	31.6	41.5	41.9	43-7	44.1	44-9	45.0	48.2	
44.5	31.9	41.8	42.2	44.0	44-3	45.I	45-3	48.4	
44.0	32.2	42.0	42.4	44.2	44.6	45-5	45.6	48.7	
43.5	32.4	42.3	42.6	44-5	44.8	45-7	45.9	40.0	1
43.0	32.7	42.6	42.9	44.8	45.I	46.0	46. ī	49.3	ŀ
42.5	52.9	42.9	43.2	45.0	45-4	46.3	46.4	49.6	
42.0	33-2	43.I	43-5	45-3	45.6	46.5	46.7	49-9	
-			43.7	45.6	45.8	46.8	47.0	50.1	1
41.5	33-5	43-4			46.1	47.0		50.4	l
41.0	33-7	43.7	44.0	45.9 46.1	46.3		47.3		1
40.5	34.0	43.9	44.2	• .	46.6	47-3	47.6	50.7	51.
40.0	34-3	44.2	44-5	46.4	40.0	47.6	47.8	51.0	31.
39 - 5	34-5	44-5		46.6	46.8	47.8	48.1	51.3	51.
39.0	34.8	44.8		46.8	47.1	48.1	48.4	51.6	52.
38.5	35.0	45.0		47-1	47-4	48.4	48.7	51.9	52.
38.0	35 - 3	45-3		47-4	47.6	48.6	48.9	52.1	52-
37-5	35-5	45.6	• • • • • • •	47.6	47.8	48.9	49.2	52-4	52.
37.0	35.8	45.9		47-9	48.1	49.2	49 - 5	52.7	53-
36.5	36.1	46.1		48.2	48.3	49.4	49.8	53.0	53-
36.0	36.3	46.4		48.5	48.6	49.7	50.0	53-3	53-
35-5	36.6	46.7		48.7	48.8	50.0	50.3	53.6	54 -
35.0	36.9	47.0		49.0	49-1	50.2	50.6	53-9	54-
34-5	37-1	47.2					50.9	54.2	54-
34.0	37-4	47-5					51.2	54-4	54-
33-5	37.6	47.8					51-5	54-7	55-
33.0	37-9	48.1	l				51.7	55.0	55-
32 5	38.1	48.3					52.0	55-3	55.
32.0	38.4	48.6					52-3	55.6	55
31.5	38.6	48.9					52.6	55.9	56.
31.0	38.9	49.2				1 1	52.8	56.1	56.
30.5	39.2	49.5					53.1	56.4	56.
30.0	39-5	49.8					53-4	56.7	57.
29.5	39-7	50.0					53-7	57.0	57.
29.5 20.0	40.0	50.3					53.9	57-3	57
28.5	40.3	50.5					54.1	57.6	57
28.0	40.5	50.8					54.4	57.8	58.
27-5	40.8	51.1	ł				54-7	58.1	58.
	41.0	51.4					55.0	58.4	58.
27.0		51.4					55.2	58.7	58.
26.5	41.3	1 -					55-5	59.0	59.
26.0	41.5	51.9					65.8	59.3	59
25-5	41.8	52.2					66.1	59.6	59
25.0	42.0	52.5					00.1	1 33.3	1 24.

^{*} Butter readings by Zeiss.
† Lard readings by Hefelmann.

CALCULATED READINGS-(Continued).

Temp. C.	Olive Oil.	Peanut Oil.	Cotton- seed Oil.	Rape- seed Oil.	Sesame Oil.	Yellow Mustard Oil.	Black Mustard Oil.	Sun- flower Oil.	Corn Oil.	Poppy- seed Oil.
35.0	57-0	59.8	61.8	62.1	62.3	63.0	64.2	64.5	65.0	C.5.5
34-5	57-2	60.0	62.1	62.4	62.5	63.3	64.5	64.8	65.3	65.8
34.0	57-4	60.3	62.3	62.7	62.8	63.6	64.8	65.1	65.6	66.1
33-5	57.7	60.6	62.5	63.0	63.1	63.9	65.1	.65.4	65.9	66.4
33.0	58.0	60.9	62.8	63.3	63.4	64.1	65.3	65.7	66.2	66.7
32.5	58.3	61.1	63.0	63.6	63.7	64.4	65.6	66.o	66.5	67.0
32.0	58.5	61.4	63.2	63.8	64.0	64.7	65.9	66.3	66.8	67.3
31.5	59.0	61.7	63.6	64.1	64.3	65.0	66.2	66.6	67.1	67.6
31.0	59.2	62.0	64.0	64.4	64.6	65.3	66.5	66.9	67.4	67.9
30.5	59-5	62.2	64.2	64.7	64.9	65.6	66.8	67.2	67.7	68.2
30.0	59-9	62.5	64.5	65.0	65.1	65.8	67.0	67.5	68.0	68.5
29.5	60.1	62.8	64.9	65.3	65.4	66.1	67.3	47-7	68.2	68.7
29.0	60.3	63.1	65.1	65.6	65.7	66.4	67.6	68.0	68.5	69.0
28.5	60.6	63.3	65.3	65.9	66.0	66.7	67.9	68.3	68.8	69.3
28.0	60.9	63.6	65.7	66.1	66.2	66.9	68.1	68.6	69.1	69.6
27-5	61.1	63.9	66.0	66.4	66.5	67.2	68.4	68.9	69.4	69.9
27.0	61.5	64.2	66.5	66.7	66.8	67.5	68.7	69.2	69.7	70.2
26.5	62.0	64.4	67.0	67.0	67.1	67.8	69.0	69.5	70.0	70.5
26.0	62.2	64.7	67.3	67.3	67.0	68.0	69.2	69.8	70-3	70.8
25.5	62.4	65.0	67.5	67.6	67.7	68.3	69.5	70. I	70.6	71.1
25.0	63.0	65.3	67.9	67.8	67.9	68.6	69.8	70-4	70.9	71.4
24.5	63.3	65.5	68.2	68.1	68.2	68.9	70.1	70.7	71.2	71.7
24.0	63.6	65.8	68.5	68.4	68.5	69.2	70-4	71.0	71.5	72.0
23.5	63.9	66.1	68.8	68.7	68.8	69.5	70-7	71.3	71.8	72.3
23.0	64.2	66.4	69.1	69.0	69.1	69.7	70.9	71.6	72.I	72.6
22.5	64.5	66.6	69.4	69.3	69.4	70.0	71.2	71.9	72-4	72.9
22.0	64.8	66.9	69.7	69.7	69.7	70.3	71.5	72.2	72.7	73.2
21.5	65.1	67.1	70.0	70.0	70.0	70.6	71.8	72-5	73.0	73-5
21.0	65.4	67.4	70.3	70.3	70.3	70.9	72.1	72.8	73 · 3	73.8
20.5	65.7	67.7	70.6	70.6	70.5	71.2	72-4	73.1	73.6	74-1
20.0	66.0	68.0	70.9	70.8	70.8	71.4	72.6	73-4	73-9	74-4
19.5	66.3	68.2	71.2	71.1	71.1	71.7	72.9	73.6	74-1	74.6
19.0	66.6	68.5	71.5	71.4	71.4	72.0	73.2	73-9	74-4	74-9
18.5	66.9	68.8	71.8	71-7	71.7	72.3	73-5	74.2	74-7	75-2
18.0	67.2	69.1	72.1	72.0	72.0	72.6	73.8	74-5	75.0	75-5
17-5	67.5	69.3	72-4	72-3	72-3	72.9	74-1	74.8	75-3	75.8
17.0	67.8	69.6	72-7	72.6	72.5	73-I	74-3	75-1	75.6	76.1
16.5	68.1	69.9	73.0	72.9	72.8	73-4	74.6	75-4	75.9	76.4
16.0	68.4	70.2	73-3	73-2	73.1	73-7	74-9	75.7	76.2	76.7
15.5	68.7	70.5	73.6	73.5	73-4	74.0	75-2	76.0	76.5	77.0
15.0	68.9	70.8	73.8	73.8	73-7	74-3	75-5	76.3	76.8	77-3

Melting-point.—A piece of small glass tubing is drawn out to a capillary open at both ends, and this is inserted into a beaker of the fat, melted at a temperature slightly above its fusing-point. A portion of the melted fat being drawn up into the capillary, the latter is removed and the fat allowed to solidify spontaneously. After an interval of not less than twelve hours, the capillary is attached by a rubber band to the stem of a delicate thermometer (preferably capable of being read to tenths of a degree), the portion of solidified fat being opposite the thermometer bulb. A test-tube containing water is held in the neck of a flask in such a manner as to be immersed in water contained in the flask, as shown in Fig. 93, the flask being held on the ring of a stand, with wire gauge interposed between flask and flame. The thermometer with attached capillary is then held immersed in the water of the test-tube and below the

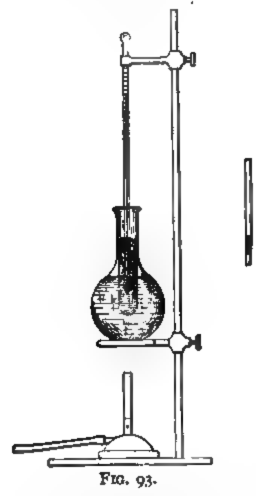


FIG. 94.

Fig. 93,—Apparatus for Determining Melting-point. Capillary tube with enclosed fat shown on the right, enlarged.

Fig. 94.—Reichert Flask with Card Inserted for Quick Evaporation.

level of the water in the flask, as shown. The water in the flask is then heated very gradually, so that the rise of temperature as shown by the thermometer does not exceed 0.5° C. per minute, the exact temperature at which fusion of the fat occurs being recorded as the melting-point.

The flame is then removed, and the temperature at which the fat solidifies is noted as the solidifying-point.

The mean of two or three determinations is usually taken as the true melting and solidifying-points.

Reichert-Meissl Process for Volatile Fatty Acids.—This process has undergone various modifications from time to time. Reichert originally used 2.5 grams of fat, but Meissl, who improved the process, used

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Fro. 95.—Apparatus for Reichert-Meissl and Polenske Distillation.

5 grams, so that the Reichert-Meissl number is now expressed on the basis of 5 grams of fat. The method is conveniently carried out as follows:

Five grams of the fat are transferred to a dry, clean Erlenmeyer flask of about 300 cc. capacity, 10 cc. of 95% alcohol are added, and 2 cc. of sodium hydroxide solution (prepared by dissolving 100 grams of sodium hydroxide in 100 cc. of water). The flask with its contents is then heated

on a water-bath with a funnel in the neck, which satisfactorily replaces the return-flow condenser originally prescribed. The heating is continued with occasional shaking till saponification is complete. This stage of the process is indicated by the appearance of the solution, which is then perfectly clear and free from fat globules.

The condenser-funnel being removed, the contents of the flask are next evaporated by continued heating over the bath to dryness. This may be hastened by inserting a card in the neck of the flask, as shown in Fig. 94, thus starting a circulatory movement to the air through the flask.

The dry soap thus formed is then dissolved by warming on the waterbath with 135 cc. of added water, shaking the flask occasionally. After cooling, 5 cc. of dilute sulphuric acid (200 parts sulphuric acid in 1 liter of water) are added, and the fatty acid emulsion formed is melted by heating the flask on the water-bath, the flask being corked during the heating. The fatty acids are completely melted when they form an oily layer on the surface of the solution.

Scraps of pumice stone joined by platinum wires are next placed in the flask to prevent bumping, and the flask is properly connected with the condenser for distilling, as shown in Fig. 95. A flask graduated at 110 cc. is used as a receiver, the funnel placed therein being provided with a loose tuft of absorbent cotton to serve as a filter. The distillation is conducted by so grading the heat that the receiving flask is filled with the distillate in about thirty minutes.

Finally the entire distillate is titrated with decinormal sodium hydroxide, using 0.5 cc. of a solution of phenolphthalein as an indicator. The number of cubic centimeters of decinormal alkali required to neutralize the acidity of the distillate from 5 grams of the fat in the manner described expresses what is known as the Reichert-Meissl number.

Lessmann and Beam's Modification.*—Five grams of the fat placed in the flask are treated with 20 cc. of a solution of soda in glycerin (20 cc. of a 50% solution of sodium hydroxide in 180 cc. of glycerin), heating the flask till the contents are completely saponified. The solution becomes perfectly clear, showing complete saponification in about five minutes, after which 135 cc. of water are added to the clear soap solution, at first drop by drop to prevent foaming; 5 cc. of the dilute sulphuric acid are then added, and the distillation conducted at once without first melting the fatty acids.

^{*} Leffmann and Beam, Select Methods of Food Analysis, p. 146.

NUI	MBER.		
•	Lowest.	Highest.	Average.
Butter	24.5	32	28.25
Cocoanut oil	24.5 6.65	7.8	7.2
Cocoa butter	0.20	0.80	0.5
Corn oil.	1.32	5.0	3.16
Lard	••••		1.10
Cottonseed oil	••••		0.95
Sesame oil	0.70	1.20	0.95
Rape oil	0.58	0.90	0.74
Olive oil	••••		0.60
Beef tallow.			0.5

EDIBLE OILS AND FATS IN THE ORDER OF THEIR REICHERT-MEISSL.
NUMBER.

Polenske Number.*—This number represents the volatile fatty acids insoluble in water, and is of value in detecting cocoanut oil in butter and other fats. The details of apparatus and manipulation here described should be closely adhered to in order to secure comparable results. Both the Reichert-Meissl and the Polenske number may be determined in one weighed portion of the fat.

Place 5 grams of the clear filtrated fat in a 300-cc. Jena flask, add 20 grams of glycerine and 2 cc. of a 50% solution of sodium hydroxide. Heat the flask on a wire gauze until the contents are completely saponified, which requires about 5 minutes, and is indicated by the clearing up of the liquid. While still hot add 90 cc. of boiled water, at first drop by drop to prevent foaming, and shake until the soap is dissolved. The solution should be completely clear and almost colorless. Rancid or oxidized fats that yield a brown soap solution should not be examined.

To the soap solution, warmed to 50°, add 50 cc. of dilute sulphuric acid (25 cc.: 1 liter) and 0.5 gram of granulated pumice stone with grains 1 mm. in diameter, then connect with the distilling apparatus shown in Fig. 95. Distil over a 0.5 mm. mesh copper gauze, using a Bunsen flame so regulated as to give a distillate of 110 cc. in 19-20 minutes, and a stream of water that will cool the distillate to about 20-23°. The room should have a temperature of about 18-22°. As soon as 110 cc. have come over, replace the flask by a 25-cc. measuring cylinder.

Without mixing the distillate place the flask for 10 minutes in water at 15°, so that the 110 cc. mark is about 3 cm. below the surface of the water. After the first five minutes, gently move the neck of the flask in

^{*} Polenske, Zeits. Unters. Nahr. Genuss., 7, 1904, p. 274. Fritsche, ibid., p. 193.

the water so that the fatty acids floating on the surface come in contact with the glass, noting at the end of 10 minutes the condition of these acids. If the butter is pure, the floating acids are either solid or form a half solid turbid mass, according as the Reichert-Meissl number is high or low; if it is adulterated with 10% or more of cocoanut oil, they form transparent oil drops. Stopper the 110-cc. flask, mix by inverting 4 or 5 times, avoiding violent shaking, filter through an 8-cm. dry filter fitted close to the funnel, and titrate 100 cc. of the liquid with tenth-normal barium hydroxide solution, thus obtaining the Reichert-Meissl number.

After the last drop of distillate has passed through the filter, wash with three 15 cc. portions of water, each of which has previously been used to rinse the condenser tube, the measuring cylinder and the 110 cc. flask. Then repeat this treatment, using 15 cc. portions of neutral 90% alcohol. Titrate the united alcoholic washings with tenth-normal barium hydroxide solution, using phenolphtalein as indicator. The number of cc. required is the Polenske number.

The following results illustrate the value of the method:

	Reichert-Meissl Number.	Polenske Number.
31 samples of butter (Polenske)	. 23.3-30.1	1.5-3.0
4 samples of cocoanut oil (Polenske)	. 6.8-7.7	16.8-17.8
Oleomargarine (Arnold)	. 0.5	0.53
Lard (Arnold)	. 0.35	0.5
Tallow (Arnold)	. 0.55	0.56

Determination of Soluble and Insoluble Fatty Acids.—A. O. A. C. Method.*—Soluble Acids.—Five grams are weighed out and transferred to an Erlenmeyer flask of the same size and in the same manner as that used for the Reichert-Meissl process. 50 cc. of alcoholic potash solution are added (40 grams of potassium hydroxide in 1 liter of 95% redistilled alcohol) and the flask, provided with a return-flow condenser, is heated on the water-bath till saponification is complete, as evidenced by the clear solution free from fat globules. The alcoholic solution of potash is preferably measured from a pipette, from which it is allowed to drain for a noted interval of time, say thirty seconds.

After complete saponification, the condenser is removed and the alcohol is evaporated by further heating. One or more blanks are pre-

^{*} U. S. Dept. of Agric., Div. of Chem., Bul. 46, p. 47; Bul. 107 (rev.), p. 138.

pared at the same time, using the same 50-cc. pipette for measuring, and applying the same time limit for draining the pipette. The blanks are first titrated, after evaporation, with half-normal hydrochloric acid, using phenolphthalein as an indicator. Then add to the flask containing the fatty acids I cc. more of the half-normal acid than is found necessary to neutralize the alkali in the blanks, after which the flask is again heated with a funnel in the neck till the fatty acids have completely separated in a layer on top of the solution. Then cool the flask in ice water till the fatty acids are solidified, after which decant the liquid portion through a filter, previously dried in the oven and weighed, into a liter flask, keeping the solid mass of fatty acids intact. Next add 200 or 300 cc. of hot water to the flask containing the fatty acids, and again melt over the water-bath till they collect as before on top, having again inserted the funnel to act as a condenser, and occasionally shaking the contents of the flask during heating. Cool as before in ice water, after which again decant the liquid from the solid mass through the same filter into the liter flask. Repeat this process of washing, melting, cooling, and decanting three times, receiving all the wash water through the same filter in the same flask. Make up the washings with water to the liter mark, and, after mixing, two portions of 100 cc. each are titrated with tenth-normal sodium hydroxide, using phenolphthalein for an indicator. Each reading is multiplied by ten to represent the total volume, and the figure thus obtained represents the number of cubic centimeters of tenthnormal alkali necessary to neutralize the acidity of the soluble fatty acids. together with the excess of half-normal acid used, amounting to 1 cc. This I cc. of half-normal acid corresponds to 5 cc. of tenth-normal alkali, hence 5 cc. are to be deducted from the total number of cubic centimeters required for the titration, the corrected figure thus obtained being multiplied by the factor 0.0088, which gives the weight of soluble fat acids in the 5 grams of the sample, calculated as butyric acid.

Insoluble Acids.—Transfer the fatty acids left in a cake in the flask from the separation of the soluble acids, to a weighed glass evaporating dish, using strong alcohol to wash them out thoroughly. Dry the filter used in the separation, transfer it to an Erlenmeyer flask, and thoroughly wash it with strong alcohol, transferring all the washings to the dish. The alcohol is then evaporated by placing the dish on the water-bath, after which it is dried for two hours in the air-oven at 100°, cooled in the desiccator, and weighed. After once heating for two hours, cooling and weighing, heat again for half an hour, cool, and weigh. If a considerable loss

in weight is found, heat for an additional half-hour. It is best, however, to avoid too prolonged heating, lest oxidation of the fatty acids should produce an increase in weight.

Hehner's Method.—Transfer the fatty acids left in the original Erlenmeyer flask to the thoroughly wet, tared filter, washing out the flask with hot water, thus bringing all the fatty acids upon the filter, which, if of good quality and thoroughly wet beforehand, will retain them. If, however, oily particles are noticed in the filtrate, they may be solidified by cooling in ice water, and afterwards removed by a glass rod and transferred to the filter. After draining dry, the funnel is immersed in cold water to solidify the fatty acids, and the filter containing them is transferred to a weighed dish, which is dried for two hours in the oven at 100°, cooled in the desiccator, and weighed, subtracting the weight of the dish and filter.

EDIBLE OILS AND FATS ARRANGED IN ORDER OF INSOLUBLE FATTY ACIDS.

Mustard oil.	96.2 to 95.1
Cottonseed oil	96 '' 95
Corn oil.	96''93
Lard	96 "93
Peanut oil	95.8
Sesame oil	95-7
Beef tallow.	95.6
Mutton tallow	95-5
Poppyseed oil	95.2 " 94.9
Rape oil	95.1
Sunflower oil	95
Olive oil	95
Cocoa butter.	94.6
Cocoanut oil	90 " 88.6
Butter	89.8 " 86.5

Saponification Number. — Koettstorjer's Method. — By the saponification number is meant the number of milligrams of potassium hydroxide necessary to completely saponify I gram of the fat. Between I and 2 grams of the fat are transferred in the usual manner (see p. 474) to an Erlenmeyer flask, and 25 cc. of the alcoholic potash solution (40 grams of potassium hydroxide free from carbonates in I liter of 95% alcohol redistilled after standing for some time with potassium hydroxide) are added with a graduated pipette, which is allowed to drain for a noted period of time, say thirty seconds. The determination should preferably

be made in duplicate. Conduct the saponification as in the case of the soluble fatty acids by heating on the water-bath. After saponification, remove from the bath, cool, and titrate with half-normal hydrochloric acid, using phenolphthalein as an indicator. Titrate also several blanks in which 25 cc. of the alcoholic potash solution are measured out with the same pipette as before, and allow to drain for the same amount of time. Subtract the number of cubic centimeters of half-normal acid necessary to neutralize the alkali in the case of the saponified fat from that necessary to neutralize the blank, multiply the result by 28.06, and divide the product by the number of grams of fat taken.

EDIBLE OILS AND FATS ARRANGED IN ORDER OF THEIR SAPONIFICATION NUMBER.

	Minimum.	Maximum.	Mean.
Cocoanut oil	246.2	268.4	257-3
Butter	225	230	227.5
Cocoa butter	192	202	197
Beef tallow	193.2	200	196.6
Lard	195.3	196.6	196
Lard oil	195	196	195.9
Cottonseed stearin	194.6	195.1	194.8
Poppyseed oil	190	198	194
Cottonseed oil	191	196.6	193.8
Peanut oil	190	197	193.5
Sunflower oil	193	194	193.5
Sesame oil	187.6	192.4	192.6
Olive oil	185	196	191.5
Corn oil.	18Š	193.4	190.
Rape oil	170.2	179.2	174.0
Black mustard oil	174	174.6	174.3
White mustard oil	170.3	174.6	172.4

The Iodine Absorption Number.—This determination is based on the well-known property of the unsaturated fatty acids to absorb a fixed amount of iodine under given conditions of time, strength of reagent, etc. Htibl's Method.*—The following reagents are necessary:

- (1) Iodine Solution, made by dissolving 26 grams of pure iodine in 500 cc. of 95% alcohol, and, separately, 30 grams of mercuric chloride in 500 cc. of the same strength of alcohol. Filter the latter solution, if necessary, and mix the two together, allowing the mixture to stand at least twelve hours before using.
- (2) Decinormal Thiosulphate Solution, made by dissolving 24.6 grams of the freshly powdered, chemically pure salt in water, and making up to 1 liter.

^{*}A. O. A. C. Method, U. S. Dept. of Agric., Bur. of Chem., Bul. 65, p. 24; Bul. 107 (rev.), p. 136.

- (3) Starch paste, prepared by boiling 1 gram of starch in 200 cc. of water for ten minutes, then cooling.
- (4) Potassium Iodide Solution, made by dissolving 150 grams of the salt in water, and making up the volume to 1 liter.
- (5) Potassium Bichromate Solution for standardizing the thiosulphate, made by dissolving 3.874 grams of chemically pure potassium bichromate in distilled water, and making up the volume to 1 liter.

The sodium thiosulphate solution is standardized as follows: 20 cc. of the potassium bichromate solution are introduced into a glass-stoppered flask together with 10 cc. of potassium iodide and 5 cc. of strong hydrochloric acid. Then slowly add from a burette the sodium thiosulphate solution, till the yellow color of the solution has nearly disappeared, after which a little of the starch paste is added, and the titration carefully continued to just the point of disappearance of the blue color. The reaction which takes place is as follows:

$$K_2Cr_2O_7 + 14HCl + 6KI = 2CrCl_2 + 8KCl + 6I + 7H_2O_2$$

The equivalent of I gram of iodine in terms of the thiosulphate solution is found by multiplying the number of cubic centimeters of the latter solution required for the above titration by 5.

If, for example, 16.4 cc. of the thiosulphate solution are required for 20 cc. of the bichromate solution, then 1 gram of iodine is equivalent to $16.4 \times 5 = 82.0$ cc. of sodium thiosulphate solution, or 1 cc. of the thiosulphate solution $=\frac{1}{8.2} = 0.0122$ gram of iodine. 1 cc. of exactly decinormal thiosulphate is theoretically equivalent to 0.0127 gram of iodine.

The thiosulphate solution may also be standardized by means of iodine. A short tube closed at one end is tared, together with another tube of such a size as to fit over the first. Into the inner tube are introduced about 0.2 gram of resublimed iodine and the tube heated until the iodine melts, after which it is closed by the second tube and the whole cooled in a desiccator and weighed. The iodine is dissolved in 10 cc. of 10% potassium iodide solution, the solution diluted with water, and the thiosulphate solution added with constant stirring until only a yellow color remains. Starch paste is then added, and the titration continued until the blue color disappears.

Method of Procedure.—Place I gram of the solid fat, or from 0.2 to 0.4 gram of oil, in a glass-stoppered flask or bottle of 300 cc. capacity.

In the case of oils, this may conveniently be done by difference, weighing first a small quantity of the oil in a beaker with a short piece of glass tubing to serve as a pipette, transferring a number of drops of the oil from the beaker to the bottle, and again weighing the beaker and contents. The number of drops of oil required for the desired weight is first ascertained experimentally.

The material may also be conveniently and accurately weighed in small, flat bottomed cylinders of glass about 10 mm. in diameter and 15 mm. high, which may be made by cutting off so-called "shell vials." Fats are introduced while melted, the weight being taken after cooling. The cylinder and fat are transferred together by means of forceps to the glass-stoppered bottle.

Dissolve the oil in 10 cc. of chloroform, and after solution has taken place, add 30 cc. of the iodine solution, shake, and set the flask in a dark place for three hours, shaking occasionally. When ready for the titration, add 20 cc. of the potassium iodide solution (the purpose of which is to keep in solution the mercuric iodide formed, which would otherwise precipitate on dilution) and 100 cc. of distilled water. Titrate the excess of iodine by the thiosulphate solution, which is slowly added from a burette till the yellow color has nearly disappeared, then add a little starch paste, and finally thiosulphate solution drop by drop until the blue color of the iodized starch is dispelled. Near the end of the reaction the flask should be stoppered and vigorously shaken, in order that all the iodine may be taken up, and sufficient thiosulphate should be added to prevent a reappearance of any blue color in five minutes.

Two blanks are conducted at the same time and in similar flasks or bottles, in exactly the same manner as in the case of the above titration, except that the fat is omitted. This is to get the true value of the iodine solution in terms of the thiosulphate solution.

Suppose, for example, in the case of the blanks, 30 cc. of the iodine solution required in one instance 46.1 cc. of sodium thiosulphate solution, and in the other 46.5 cc. The mean is 46.3. Suppose 15.1 cc. of thiosulphate solution were required for the excess of iodine remaining over and above that absorbed by 1 gram of the fat in the above process. Then the thiosulphate equivalent to the iodine absorbed by the fat would be 46.3-15.1=31.2 cc., and the per cent of iodine absorbed would be $31.2\times0.0122\times100=38.06$.

EDIBLE OILS AND FATS ARRANGED IN ORDER OF THEIR HÜBL NUMBER.

	Lowest.	Highest.	Average.
Poppyseed oil	132.6	143-3	138
Sunflower oil	118	133.3	125.7
Corn oil	. 111.2	130	120.6
Cottonseed oil	108	110	100.5
Sesame oil	103	115	100
Rape oil	94	105	99.5
Black mustard oil	9 6	110	103
White mustard oil	92.I	97.7	94.9
Peanut oil	8 3	103	93
Cottonseed stearin		103.8	QI.2
Olive oil		88	83.5
Lard oil		85	70.5
Lard		70	58.
Beef tallow		47-5	41.4
Mutton tallow		46.2	39-5
Cocoa butter		41.7	34.9
Butter		37.9	33-3
Cocoanut oil		9.5	8.7

The Hübl method has long been almost universally used for estimating the per cent of iodine absorbed, but is open to serious objections, chief of which are the tendency of the iodine solution to lose strength, and the length of time required to insure saturation of the oil with the iodine.

Of late two other methods have come into prominence, viz., the Wijs and the Hanus. The reagents in both these methods hold their strength for months without change, and the time required for carrying out the reaction in the case of most of the edible oils and fats is very short.

Of the three methods, that of Hanus has the advantage of greatest simplicity in the composition and preparation of the chief reagent.

Tolman and Munson* have shown that with oils and fats having iodine numbers below 100, the three methods give practically identical figures, while with oils having high iodine numbers, the Wijs and Hanus methods give higher results than the Hübl, but are doubtless more nearly correct.

The following are comparative results of the three methods:*

^{*} Jour. Am. Chem. Soc., 25 (1903), p, 244.

Number of Analyses.			Hūbl's Number (3 bours).	Wijs's Number (30 min.).	Hanus' Number (30 min.).	Difference between Wijs and Hubl.	Difference between Hanus and Hübl.
1	Cocoanut oil		8.93	9.05	8.60	+0.12	-0.33
2	Butter-	minimum	34.8	35.9	35-4	+1.1	+0.6
		maximum	35-3	36.2	35 · 3	+0.9	+0.0
I	Oleo oil		42.6	43-5	43-3	+0.9	+0.7
4	Oleomargarine	-minimum	52.5	52.9	52.0	+0.4	-0.5
	1	maximum	66.3	66.0	64.8	-0.3	-1.5
2	Lard oil-	minimum	69.3	70.5	69.8	+1.2	+0.5
46	Olima all	maximum	73-7	74.5	73-9	+0.7	+0.2
3 6	Olive oil—	minimum	79.2	79-9	80.6	+0.7	+1.4
		maximum	89.8 84.0	91.4	90.0 84.6	+1.6	+0.2
3	Peanut oil-	average minimum	94.5	85.3 95.2	94.I	+1.3	-0.I
3	2 condition	maximum	107.7	100.5	107.7	+1.8	+0.0
5	Mustard oil-	minimum	98.4	104.3	103.8	+5.9	+5.4
•		maximum	113.0	118.2	116.8	+5.2	+3.8
2	Rape oil-	minimum	100.2	104.1	102.8	+3.9	+2.6
	i -	maximum	101.3	105.7	105.2	+4.4	+3.8
1	Sunflower oil .		106.4	109.2	107.2	+2.8	+0.8
3	Cottonseed oil-		103.8	105.3	105.2	+1.5	+1.4
		maximum	106.2	107.3	107.8	+1.1	+1.6
I	Sesame oil	• • • • • • • • • • • • • • • • • • • •	106.4	107.0	106.5	+0.6	+0.1
3	Corn oil—	minimum	119.0	122.2	119.6	+3.0	+0.4
2	Poppygood all	maximum	123.3	129.2	126.0	+5.8	+2.7
2	Poppyseed oil—		133.4	135.2	132.9	+1.8	-0.5
	1	maximum	134.9	139.1	138.4	+4.2	+3-5
				<u> </u>		<u> </u>	<u> </u>

Hanus' Method.*—Reagents.—Iodine Solution.—Dissolve 13.2 grams of pure iodine in 1 liter of pure glacial acetic acid (99%), and to the cold solution add 3 cc. of bromine, or sufficient to practically double the halogen content when titrated against the thiosulphate solution, but with the iodine slightly in excess.

Decinormal Thiosulphate Solution, Starch Paste, and Potassium Iodide Solution, as in Hübl's method.

Method of Procedure.—Proceed as in Hübl's method, substituting 30 cc. of the Hanus iodine reagent for that of Hübl, stirring the solution before adding the water, and, instead of adding 20 cc. of the potassium iodide solution, use only 10 cc.

^{*}Zeits. f. Unters. Nahr. u. Genuss., 4 (1901), p. 913. Also Hunt, Jour. Soc. Chem Ind. 21 (1902), p. 454; U. S. Dept. of Agric., Bur. of Chem., Bul. 107 (rev.), p. 136.

Only half an hour is required for full saturation of the oil by the iodine in the Hanus method, as against three hours in the Hübl. In case of the non-drying oils and fats, the reaction takes place in from eight to fifteen minutes, though it is best to let the flask set for half an hour at least, in all cases. With oils having an iodine number in excess of 100, Tolman and Munson recommend one hour's standing.

On account of the high coefficient of expansion of acetic acid, care should be taken that the temperature is the same when the iodine solution is measured for the blank and for the determination, as otherwise a serious error may be introduced.

Wijs's Method.*—Reagents.—Iodine Solution.—Dissolve 13.2 grams of pure iodine in 1 liter of pure glacial acetic acid, and pass through the larger portion of this solution a current of carefully washed and dried chlorine gas † until the solution is practically decolorized. Finally add enough of the original solution of iodine in acetic acid to restore the iodine color, so that there is a slight excess of iodine.

Hunt's Modified Iodine Solution.—Dissolve 10 grams of iodine trichloride in 1 liter of pure glacial acetic acid, and finally add and dissolve 10.8 grams of pure iodine.

Other Reagents, as in the Hübl and Hanus methods.

Method of Procedure.—Proceed as in the Hanus method, observing the same precautions, the only difference being in the use of the Wijs iodine reagent.

Wijs recommends the following periods of time for absorption of the iodine in the case of various oils: For non-drying oils and fats, such as peanut and olive oil,‡ fifteen minutes; for semi-drying oils, such as cottonseed, rape, sesame, corn, and mustard, thirty minutes; for drying oils, such as sunflower and poppyseed, one hour.

The Bromine Index or Bromine Absorption Number.—The measure of the amount of bromine absorbed by the oils and fats is a useful factor. By the bromine index is understood the weight of bromine which is

^{*} Ber. d. chem. Ges., 31 (1898), p. 750.

[†] The chiorine is conveniently prepared by treatment of bleaching powder with dilute sulphuric acid, using gentle heat, and washing the gas by passing through strong sulphuric acid.

[‡] For butter, oleo oil, lard oil, and cocoanut oil, fifteen minutes is sufficient.

absorbed by I gram of a given oil. The bromine index of various oils has been determined as follows:

	Bromine Index.	Observer.
Poppyseed. Mustard. Sesame. Cottonseed Rape Peanut. Olive.	0.763 0.695 0.645 0.632 0.530	Levallois Girard Levallois " Girard Levallois

Method of Levallois.—Five grams of the oil are saponified with alcoholic potash in a 50-cc. graduated flask by the aid of a gentle heat. At the end of the saponification and after cooling, the flask is filled to the mark with alcohol, and, after shaking, 5 cc. are removed by means of a pipette and transferred to a flask. A slight excess of hydrochloric acid is added to set free the fatty acids, and from a burette a standardized solution of bromine water is run in till with constant shaking a permanent yellow color persists. The bromine is previously standardized with potassium iodide and sodium thiosulphate. The weight of bromine fixed by 1 gram of the fat is then calculated.

Mill's Method.—Modified.†—Dissolve o.1 gram of the filtered and dried fat in 50 cc. of carbon tetrachloride or chloroform in a 100-cc. stoppered bottle. From a burette a standard solution of bromine in carbon tetrachloride, approximately tenth-normal (8 grams to a liter), is slowly added to the oil solution till, after fifteen minutes, a permanent coloration remains. The amount of bromine absorbed is calculated by comparing with the color similarly produced in a blank experiment, or an excess of bromine solution may be run in and the solution titrated back with a standard solution of thiosulphate, using potassium iodide and starch.

Thermal Tests.—The rise in temperature produced by the action of certain reagents on various oils and fats, when applied in a definite manner, has been found to be of considerable value, especially in the case of sulphuric acid and of bromine.

^{*} Villiers et Collin, Les Substances Alimentaires, p, 680.

[†] Allen, Commercial Org. Analysis, II, part 1, p. 63.

The Maumené Test,* or thermal reaction with sulphuric acid, is most readily carried out in a beaker of say 150 cc. capacity, which is set into a larger beaker or vessel of any kind, the space between the two being packed with felt or cotton waste. The inner beaker is removed, and into it is weighed 50 grams of the oil. It is then replaced and the packing adjusted, if necessary, after which the temperature of the oil is noted with a thermometer. From a burette containing the strongest sulphuric acid of the same temperature as the oil, 10 cc. are run into the beaker, at the same time stirring the mixture of acid and oil with the thermometer. The temperature rises somewhat rapidly, and remains for an appreciable time at its maximum point, which should be noted. The difference in degrees centigrade between the initial temperature of the oil and the maximum temperature of the mixture expresses the Maumené number.

With certain oils, as cottonseed, considerable frothing ensues when concentrated acid is employed, making an accurate determination of the Maumené number somewhat difficult. In this case it is better to employ a somewhat weaker acid, and to express results in terms of what is called the "specific temperature reaction." This is the result obtained by dividing the rise of temperature in the case of the oil by the rise of temperature in the case of water, using the same strength of acid, and multiplying the quotient by 100. Indeed, it is of importance in all cases to compare results on oils with those obtained by carrying out the same test on water.

Bromination Test.—This test depends upon the avidity with which the oils and fats absorb bromine, the rise in temperature caused by the reaction being measured in this case rather than the actual amount of bromine absorbed, as in the case of the iodine absorption. Indeed, there is such a close relation between the iodine number and the heat of bromination, that when one is determined the other may be calculated quite closely by multiplying by a factor. In view of the fact that the heat of bromination is much more readily determined than the iodine number, it is often convenient to calculate the latter from the former, the result in the case of the edible oils and fats being quite sure to fall within the limits of variation of the iodine number of different oils of the same class. The bromination test was devised by Hehner and Mitchell,† who employed a vacuum jacketed tube for a calorimeter in which to make the test. Various modifications have been suggested both in the

^{*} Maumené, Compt. Rend., XXXV (1852), p. 572.

[†] Analyst, XX (1895), p. 146.

apparatus employed and in the manner of diluting the oil and applying the reagent. The calorimeter employed by Gill and Hatch,* Fig. 96, is conveniently made and is very satisfactory. It consists of a long, narrow, flat-bottomed tube, held by a cork in a small beaker, in such a manner that it is surrounded by an air jacket. The small beaker is set into one of larger size, the space between the two being packed with cotton waste. Five grams of the oil or fat are dissolved in 25 cc.

 A_{+}

В.

F1G. q6.

A. Gill and Hatch's Calorimeter for the Bromination Test with Oils.

B. Wiley's Pipette for Measuring Bromine in Chloroform.

of chloroform or carbon tetrachloride, and 5 cc. of this solution (containing 1 gram of the oil) are transferred by a pipette to the inner tube of the above calorimeter, being careful not to let it flow down the sides of the tube. The temperature of the oil is then taken by a thermometer graduated to 0.2°. The bromine reagent, which should be freshly prepared, is made up by measuring from a burette one part by volume of bromine into four parts of chloroform or carbon tetrachloride. The reagent is transferred to a measuring-flask devised by Wiley,† consisting of a side-necked filter-flask provided with a per-

^{*} Jour. Am. Chem. Soc., XXI (1899), p. 27. Gill, Oil Analysis, p. 50.

[†] Jour. Am. Chem. Soc., XVIII (1896), p. 378.

forated rubber stopper into which the stem of a 5-cc. pipette is fitted, Fig. 96. A bulb on the side-neck serves to fill the pipette. This pipette, filled to the mark with the bromine reagent (which should be at the same temperature as the oil solution in the calorimeter), is first covered by the finger and removed, and its contents of 5 cc. allowed to flow down the sides of the inner tube of the calorimeter and mingle with the oil without stirring. The rise in temperature is very quick, and the highest point is noted. The difference between the highest and the initial temperature constitutes the heat-of-bromination number.

This number, in the case of Gill and Hatch's calorimeter, is somewhat lower than when a vacuum jacketed tube is employed, and differs somewhat with the diluent of the oil and bromine. In spite of these variations and that due to the personal equation, concordant results may be obtained with the various oils, when the method is carried out under precisely the same conditions. The analyst should carefully work out the test several times with a particular oil till the results agree, and should then with equal care determine the iodine number of the same oil. The iodine number, divided by the heat-of-bromination number, gives the factor which is to be employed under the same conditions for calculating one constant from the other. In the case of Hehner and Mitchell's work with the vacuum tube, measuring I cc. of undiluted bromine into I gram of oil dissolved in IO cc. of chloroform, it was found that the factor to be used in calculating the iodine number was 5.5.

The following are some of the results on edible oils obtained by Hehner and Mitchell:

Oil.	Heat of Bromination.	Iodine Number.	Calculated Iodine Number.
Lard	21.5	57.15 37.07 80.76 122 107.13	58.3 36.3 82.5 118.2 106.7

As in the case of the Maumené test with sulphuric acid (wherein the rise in temperature of sulphuric acid and water is taken as a standard), it is convenient to employ some standard for the bromination test, whereby varying results due to difference in apparatus, etc., may be compared.

In this case Gill and Hatch found that sublimed camphor may be prepared sufficiently pure to be used for such a standard. Applying the bromination test with their calorimeter, as above described, to 5 cc. of a

solution of 7½ grams of camphor in 25 cc. of carbon tetrachloride, an average rise in temperature of 4.2° was obtained, and the specific temperature reaction is calculated for each oil by dividing the heat of bromination by this number. Furthermore, by dividing the iodine number of several oils by this specific temperature reaction, the factor to be employed for the calculation of the iodine number was found to be 17.18, as in the following cases:*

011	Specific Tem-	Iodine N	umber.
Oil	perature Reaction.	Calculated.	Found.
Prime lard	3-705 4-096 4-762 5-667 6-381	63.8 70.3 81.8 97.3 109.5	63.8 73.9 82.0 103.0 107.8

The Acetyl Value. — On heating fats with acetic anhydride they become "acetylated"; i.e., the hydrogen atom of their alcoholic hydroxyl group is exchanged for the acetic acid radicle, in accordance, for example, with the following reaction:

$$\begin{array}{c} C_{17}H_{32}(OH)COOH + (C_2H_3O)_2O = C_{17}H_{32}(O,C_2H_3O)COOH + C_2H_4O_2. \\ \begin{array}{c} \text{Ricinoleic} \\ \text{acid} \end{array} \\ \begin{array}{c} \text{Acetic anhy-} \\ \text{dride} \end{array} \\ \end{array}$$

By the actyl value is meant the number of milligrams of potassium hydroxide necessary to neutralize the acetic acid formed by the saponification of 1 gram of the acetylated fat.

Lewkowitsch's method of procedure is as follows: 10 grams of the oil are boiled with an equal volume of acetic anhydride for two hours in a flask with a return-flow condenser, and the mixture is then transferred to a large beaker containing 500 cc. of water, and boiled for half an hour. To prevent bumping, a current of carbon dioxide is slowly passed through it during the boiling, introduced through a finely drawn, bent glass tube reaching nearly to the bottom of the beaker. The mixture on standing separates into two layers, of which the lower, or aqueous layer, is siphoned off, and the oily layer boiled with fresh portions of

^{*} Gill, Oil Analysis, p. 128.

water, which are in turn siphoned off, the operation being repeated till the wash water tests free from acid by litmus paper.

The acetylated fat is then separated from the water by drying at 100° in an oven.

From 2 to 4 grams of the acetylated fat is weighed into a flask, and saponified with alcoholic potash in precisely the same manner as for the determination of the saponification number. Evaporate the alcohol and dissolve the soap in water. One of two methods may be carried out for freeing the acetic acid for titration, one by distillation and the other by filtration.

For the former or distillation process, acidify the aqueous solution of the soap with 1:10 sulphuric acid, and distill in the same way as in the Reichert process, excepting that in this case from 600 to 700 cc. of distillate must be obtained, so that water should be added from time to time through a stoppered funnel fixed in the cork of the distilling-flask. The distillate should be received in a funnel with a loose cotton plug, so as to filter it free from insoluble acids mechanically carried over. The filtrate is titrated with tenth-normal sodium hydroxide, using phenol-phthalein as an indicator. The number of cubic centimeters of alkali used is multiplied by 5.61, and the product divided by the number of grams of acetylated fat taken. The result is the acetyl value.

If the filtration process is used (which is more rapid and should give concordant results with the distillation process), the exact amount of alcoholic potash used in the saponification should be accurately measured in carrying out the former part of the test, and the exact number of cubic centimeters of standard acid corresponding to the amount of alkali employed should be added to the aqueous soap solution. The mixture should be gently warmed, and the fatty acids will gather in a layer at the top. These are filtered off and washed, till free from acid, with boiling water. The filtrate is titrated with tenth-normal sodium hydroxide, and the acetyl value calculated as in the distillation process.

EDIBLE OILS ARRANGED IN ORDER OF ACETYL VALUE.

	Average.
Cottonseed oil	. 18.0
Rape oil	. 14.7
Poppyseed oil	. 13.1
Sesame oil	
Olive oil	. 10.6
Peanut oil	3-4

The Valenta Test.—This depends upon the solubility of the oil in glacial acetic acid. Pour from 3 to 5 cc. of the oil into a test-tube, and add an equal volume of glacial acetic acid (specific gravity 1.0562). Place a thermometer in the tube and warm gently till the oil goes into solution. Then allow the mixture to cool, and observe the temperature at which the solution begins to appear turbid.

Castor oil and oil of the olive kernel are soluble in glacial acetic acid at ordinary temperatures, while rape and mustard seed oils are insoluble even in the boiling acid.

Elaïdin Test.—This is based on the conversion by nitrous oxide of liquid olein into the solid elaïdin, a crystalline compound isomeric with olein, while other common glycerides remain liquid under treatment with this reagent. By the consistency of the final product, when subjected under certain conditions to the action of nitrous oxide, some idea as to the character of the oil may be gained.

Manipulation.—To carry out the test according to Pontet (modified), weigh 5 grams of the oil into a beaker, add 7 grams of nitric acid (specific gravity 1.34) and about 0.5 gram of copper wire. Place the beaker in water at 15° and stir thoroughly with a glass rod in such a manner as to make an intimate mixture of the oil and the evolved nitrous oxide gas. After the wire has been dissolved, add another piece of about the same size and again stir vigorously. Set aside for about two hours, at the end of which, in the case of pure olive, almond, peanut, or lard oil, it will have been changed into a solid white mass.

Nearly all the seed oils, especially cottonseed and mustard, are turned into a pasty or buttery mass.

Another modification of Pontet's test consists in mixing 10 grams of the oil, 5 grams of nitric acid (specific gravity 1.38), and 1 gram of mercury in a test-tube, shaking for three minutes and allowing to stand twenty minutes, when it is again shaken.

The behavior of various oils after that time on further standing is as follows:

	Solidif	ied after
Olive oil	60 n	ninutes
Peanut oil	80	"
Sesame oil	185	"
Rape oil		

Free Fatty Acids.*—Weigh 20 grams of the oil or fat into a 150-cc. Erlenmeyer flask, and add 50 cc. of 95% alcohol, which has previously

^{*} Allen Com. Org. Anal., 3d ed., vol. 2, pt. 1, p. 105.

been carefully neutralized with a weak solution of sodium hydroxide, using phenolphthalein as an indicator. Warm the mixture to about 60°, and add carefully from a burette tenth-normal sodium hydroxide (using the above indicator) till a pink color is produced, shaking thoroughly during the titration.

The result may be reported in terms of percentage of oleic acid (each cubic centimeter of tenth-normal alkali is equivalent to 0.0282 gram of oleic acid) or as the "acid number," by which is meant the number of cubic centimeters of tenth-normal alkali necessary to saturate the free acid in 1 gram of the fat or oil.

Constants of the Free Fatty Acids.—Often much information as to the character of an oil or fat may be obtained by determining such constants of its fatty acids as the melting- and solidifying-point, the iodine number, etc.

To prepare the fatty acids for examination, saponify a quantity of the oil or fat with alcoholic potash, evaporate the alcohol, and dissolve the soap in hot water. Decompose the soap by the addition of an excess of hydrochloric or sulphuric acid, continuing the heating till the fatty acids rise in a layer to the top of the liquid, from which they may be removed. The melting-point, iodine number, ecc., are determined as with the oil or fat itself.

Solidifying-point of the Fatty Acids, or Titer Test.—Modified Wolfbauer Method.*—Saponify 75 grams of fat in a metal dish with 60 cc. of 30% sodium hydroxide (36° Baumé) and 75 cc. of 95% by volume alcohol or 120 cc. of water. Boil to dryness, with constant stirring to prevent scorching, over a very low flame, or over an iron or asbestos plate. Dissolve the dry soap in a liter of boiling water, and if alcohol has been used, boil for forty minutes in order to remove it, adding sufficient water to replace that lost in boiling. Add 100 cc. of 30% sulphuric acid (25° Baumé) to free the fatty acids, and boil until they form a clear, transparent layer. Wash with boiling water until free from sulphuric acid, collect in a small beaker, and place on the steam bath until the water has settled and the fatty acids are clear; then decant them into a dry beaker, filter, using a hot-water funnel, and dry twenty minutes at 100° C.

When dried, cool the fatty acids to 15 or 20° C. above the expected titer, and transfer to the titer tube, which is 25 mm. in diameter and 100 mm. in length (1 by 4 inches), and made of glass about 1 mm. in thickness. Place in a 16-ounce saltmouth bottle of clear glass, about 70 mm. in diameter and 150 mm. high (2.8 by 6 inches), fitted with a cork, which is perforated so as to hold the tube rigidly when in position. Suspend the

^{*} A. O. A. C. Method, U. S. Dept. of Agric., Bur. of Chem., Bul. 107, p. 135.

thermometer, graduated to 0.10° C., so that it can be used as a stirrer, and stir the mass slowly until the mercury remains stationary for thirty seconds. Then allow the thermometer to hang quietly, with the bulb in the center of the mass, and observe the rise of the mercury. The highest point to which it rises is recorded as the titer of the fatty acids.

Test the fatty acids for complete saponification as follows:

Place 3 cc. in a test tube and add 15 cc. of alcohol (95% by volume). Bring the mixture to a boil and add an equal volume of ammonium hydroxide (0.96 sp. gr.). A clear solution should result, turbidity indicating unsaponified fat. The titer must be made at about 20° C. for all fats having a titer above 30° C. and at 10° C. below the titer for all other fats.

The thermometer must be graduated in tenth degrees from 10° to 60°, with a zero mark, and have an auxiliary reservoir at the upper end, also one between the zero mark and the 10° mark. The cavity in the capillary tube between the zero mark and the 10° mark must be at least 1 cm. below the 10° mark, the 10° mark to be about 3 or 4 cm. above the bulb, the length of the thermometer being about 15 inches over all. The thermometer is annealed for 75 hours at 450° C., and the bulb is of Jena normal 16¹¹¹ glass, moderately thin, so that the thermometer will be quick acting. The bulb is about 3 cm. long and 6 mm. in diameter. The stem of the thermometer is 6 mm. in diameter and made of the best thermometer tubing, with scale etched on the stem, the graduation to be clear cut and distinct, but quite fine.*

Unsaponifiable Matter.—As will be seen by reference to the table on page 509, the unsaponifiable matter in pure edible oils and fats is comparatively insignificant in amount, consisting largely of cholesterol or phytosterol. A high content of unsaponifiable matter is indicative of adulteration, pointing to the presence of mineral or coal-tar oils, or to paraffin.

Determination of Unsaponifiable Matter.†—Weigh 7 to 10 grams of the fat or oil in a 250-cc. flask, and saponify by boiling with 25 cc. of alcoholic potassium hydroxide and 25 cc. of alcohol under a returnflow condenser. After saponification, add 30 to 40 cc. of water, and bring to the boiling-point. Cool and transfer the contents from the flask to a separatory funnel, washing out the flask first with a small amount

^{*} Tolman, U. S. Dept. of Agric., Bur. of Chem., Bul. 90, p. 75.

[†] Hönig and Spitz, Jour. Soc. Chem. Ind., 1891, p. 1039.

of 50% alcohol, and finally with 50 cc. of petroleum ether (B.P. 40°-70°), adding both washings to the separatory funnel. Shake the latter thoroughly, but avoid if possible forming an emulsion. If the latter persists in forming, add a volume of water equal to that of the soap solution, which will sometimes break it up. After separation of the petroleum ether layer, draw off the underlying soap solution into a beaker. and wash the petroleum ether two or three times with 50% alcohol, which is drawn off and added to the soap solution. The petroleum ether is then run into a tared Erlenmeyer flask, and the soap solution extracted twice more with fresh portions of petroleum ether, washing the ether each time with 50% alcohol as before and then transferring the ether to the tared flask. The petroleum ether is then removed by placing the flask on the water-bath, bumping being prevented by means of a spiral of platinum wire weighed with the flask. Finally remove all traces of remaining ether by blowing hot air through the flask, or, in the absence of mineral oils (some of which are volatile), dry in the water-oven to constant weight, cool in a desiccator, and weigh.

Cholesterol and Phytosterol.—These are monatomic alcohols, and combine with the fatty acids forming esters. Both respond to the same reactions, and are separated by the same process from the oils and fats in which they occur. Phytosterol was long thought to be the same as cholesterol, and some confusion seems to have arisen from the fact that early writers purport to have found cholesterol in vegetable oils, when in reality the substance was phytosterol. The latter was first distinguished from cholesterol by Hesse, who named it.

Cholesterol (C₂₆H₄₄O) crystallizes in white, nacreous, monoclinic laminæ, having a melting-point of 145° and specific gravity 1.067. Its reaction is neutral, it is devoid of taste or smell, insoluble in water, sparingly soluble in cold, but readily soluble in boiling alcohol, and soluble in ether, chloroform, methyl alcohol, benzene, and oil of turpentine. It sublimes unchanged at 200°, but at higher temperatures decomposes.

Commercial cholesterol is obtained from wool oil and is known as lanolin, being used largely in medicine as a basis for ointment.

Cholesterol occurs also in the yolk of eggs, in many animal secretions, and in most animal oils and fats.

It separates in laminated, transparent crystals from a mixture of 2 volumes alcohol and 1 volume ether, and in the form of anhydrous needles from chloroform.

Phytosterol (C26H44O,H2O) is most abundantly found in the legu-

minous seeds, and is prepared commercially from these, especially from peas and lentils. It is a constituent of most vegetable oils.

It crystallizes in slender, glittering plates from chloroform, ether, and petroleum ether, and from alcohol in tufts of needles. In solubility it much resembles cholesterol, but its melting-point from 132° to 134° is lower.

Determination of Cholesterol and Phytosterol.—Method of Forster and Reichmann.*--50 grams of the oil or fat are boiled for five minutes in a flask connected with a reflux condenser with two successive portions of 75 cc. of 95% alcohol, and in each case the alcoholic solution is separated by means of a separatory funnel. The combined alcoholic solutions are then boiled in a flask provided with a funnel in the neck, till one-fourth of the alcohol is evaporated, and then poured into an evaporating dish and brought to dryness. The residue is then extracted with ether, and the ether solution is evaporated to dryness, taken up again with ether, filtered, evaporated once more, and dissolved in hot 95% alcohol, from which it is allowed to crystallize. Cholesterol or phytosterol will crystallize out under these conditions, and may be weighed.

Distinguishing between Cholesterol and Phytosterol.—It is sometimes of importance to determine which of these substances is present in an oil, or whether indeed both occur. Confirmatory proof as to the presence of vegetable in animal oils may, for instance, be established by showing whether the unsaponifiable residue in the sample contains cholesterol or phytosterol or both. Hehner † has made use of this test in determining the presence of cottonseed oil in lard.

The most ready means of distinguishing between cholesterol and phytosterol is furnished by the marked difference between the form of the crystals, the manner of crystallization of the two substances, and the melting points of the acetates.

Separation and Crystallization of Cholesterol and Phytosterol.—Bömer's Method.‡—Saponify 100 grams of the fat by heating in a liter Erlenmeyer flask on a boiling water bath with 200 cc. of alcoholic potash solution (200 grams of potassium hydroxide+1 liter of alcohol). The flask should be provided with a perforated rubber stopper, through which passes a glass tube 700 cm. long, which serves as a reflux condenser. During the first part of the heating shake often and vigorously until the solution is clear, after which continue the heating one-half to one hour longer with occasional shaking.

^{*} Analyst, 22, 1897, p. 131.

[†] Ibid., 13, 1888, p. 165.

[‡] Zeits. Unters. Nahr. Genuss., 1, 1898, p. 31.

While still warm, transfer to a separatory funnel of about 1.5 liters capacity, rinsing the flask with 400 cc. of water. When cool, add 500 cc. of ether, shake vigorously for one-half to one minute, opening the cock repeatedly, and allow to stand for two to three minutes until the liquids separate. Remove the ether solution to a flask, and distil off the ether, using a few pieces of pumice stone to prevent bumping. Shake the soap solution two to three more times in the same manner with 200 to 250 cc. of ether, add the ether solution after each shaking to the residue in the distilling flask, and distil off the ether.

Usually a small amount of alcohol remains in the flask after removal of the ether, which may be removed by heating on a boiling water bath in a blast of air. To saponify any remaining fat, add 20 cc. of the alcoholic potash solution, and heat for five to ten minutes as before. Transfer to a small separatory funnel, rinse with 40 cc. of water, cool and shake with 150-200 cc. of ether from one-half to one minute, allow to stand two to three minutes, and draw off the lower layer. Wash the ether solution three times with 10-20 cc. of water, filter, to remove drops of water, into a small beaker, and remove the ether by cautious evaporation on the water bath, thus obtaining the crude cholesterol or phytosterol.

The unsaponifiable residue, which may be weighed after drying, in the case of animal fats shows beautiful radiating crystals, and consists largely of cholesterol, while in the case of vegetable fats it consists largely of phytosterol. Dissolve the residue in 4-20 cc. of absolute alcohol with the aid of heat, and allow to crystallize slowly in a shallow dish.

The crystallization in the case of cholesterol alone begins from the margin of the liquid and gradually extends inward toward the center, forming a uniformly bright, thin, colorless film over the whole surface. This film is best removed with a knife or spatula and pressed between filter-paper. The film will be seen, even megascopically, to be composed of large, glossy plates with a silk-like luster. After the removal of the first film a second will form similar to the first, but composed as a rule of smaller crystals. These are removed in like manner, dried between filters, and added to the first in a glass. After the second crop, the mother liquid is thrown away. The crystals are then redissolved in absolute alcohol, and again allowed to separate out, being repeatedly recrystallized till the melting-point is constant. In lard and most fats the crystals were found pure by Bömer after the second crystallization.

Phytosterol is crystallized with greater difficulty, especially when derived from seed oils, on account of the presence of pigments and other

foreign matter. The first procedure is the same as above described for cholesterol, the crystals being allowed to separate slowly out of a solution in absolute alcohol. Unlike cholesterol, no film is formed on the surface, but needles (sometimes I cm. in length) are gradually eliminated, beginning at the margin and extending inward mostly at the bottom. In concentrated solutions, fine needles would be uniformly deposited through the liquid. These are best separated from the mother liquid by filtration, as they are not easily taken out with a knife. They may be washed on the filter with small amounts of absolute alcohol for microscopical examination, or repeatedly recrystallized, as in the case of cholesterol, till the melting-point is constant.

1. Cholesterol Crystals.—When crystallized separately under above conditions, cholesterol crystals viewed under the microscope show generally rhomboidal forms of plates, as in Fig. 97, but sometimes with a reenter-

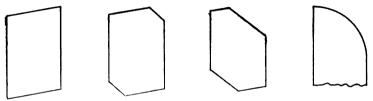


Fig. 07.—Cholesterol Crystals under the Microscope. (After Bömer.)

ing angle. The plates are often grown together in masses. The most characteristic forms are found from the first crystallization or from the first film removed. Sometimes quadrilateral crystals predominate among the plates, often also the other shapes shown are found most numerous.

2. Phytosterol Crystals.—Pure phytosterol crystallizes in needles or narrow plates, arranged commonly in star form or in bunches. The most common forms are shown in Fig. 98, best conditions as to shape of crystals being obtained from slow crystallization, in which case the needles are finer and more regular.

The crystals are commonly in the form of long, narrow plates, thin and slender, often pointed at both ends. Sometimes the points are lacking, or the ends are beveled. The more frequently they are recrystallized, the larger and more varied are the crystal forms. The broad, hexagonal and quadrilateral plates shown are products of re-

crystallization; the shorter forms are rarely met with. Sometimes various forms are found side by side in the same crystallization.

Phytosterol crystals, from a second or third recrystallization, sometimes grow together in bunches resembling at first glance to the naked eye the cholesterol masses. They never do this in the first crystallization, whereas in the case of cholesterol the growing together in masses is very characteristic of the first crystallization.

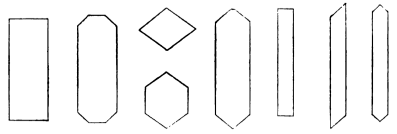


Fig. 98.—Phytosterol Crystals. (After Bömer.)

Thus for purposes of distinguishing between the two the product of the first crystallization is best observed.

3. Crystals of Mixed Cholesterol and Phytosterol.—In mixtures of the two they do not crystallize separately, but when in nearly equal proportion, or with phytosterol predominating, the crystals much resemble phytosterol. Even when cholesterol predominates to the extent of 20 parts to 1 of phytosterol, the mode of crystallization leans most toward that of phytosterol, though the needles are of different shape. Such a mixture, for instance, does not form in a film like cholesterol, but, like phytosterol, comes out in needle-like bunches. The needles, however, are more often like those shown in Fig. 99 when viewed under the micro-

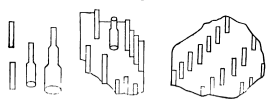


Fig. 99.—Characteristic Forms of Crystallization of Mixed Cholesterol and Phytosterol (After Bömer.)

scope, showing needles for the most part squarely cut off at the ends, and sometimes placed end to end, and of varying diameter, giving the appearance of a spy-glass. When cholesterol predominates over phytosterol 50 to 1, the plates resemble those of cholesterol.

Bömer's Phytosterol Acetate Test for Vegetable Fats.*—Dissolve the crude cholesterol or phytosterol, or the mixture of the two, obtained by Bömer's method, as described on page 503, in the smallest possible amount of absolute alcohol, and allow to crystallize. Examine under the microscope the first crystals that separate, comparing with the cuts and descriptions given in the preceding section. Remove the alcohol completely by evaporation on the water bath, add 2 to 3 cc. of acetic anhydride, cover with a watch glass, and boil for one-fourth minute on a wire gauze; then remove the watch glass, and evaporate the excess of acetic anhydride on the water bath. Heat the residue with sufficient absolute alcohol to dissolve the esters, and add enough more to prevent immediate crystallization on cooling. Cover until the room temperature is reached and allow to crystallize.

After one-half to one-third of the liquid has evaporated and the greater part of the esters have crystallized, transfer the crystals to a small filter by the aid of a small spatula, rinsing with two portions of 2 to 3 cc. of 95% alcohol. Return the crystals to the crystallizing dish, dissolve in 5 to 10 cc. of absolute alcohol, and again allow to crystallize. After the greater part of the crystals have separated, collect on a filter as before. Repeat the recrystallization several times (5 to 6 is usually sufficient), determining the melting point of the crystals after each recrystallization beginning with the third.

If after the last crystallization the corrected melting point of the crystals is above 116°, the presence of a vegetable fat or oil is indicated, if it is 117° or higher the proof may be regarded positive.

The standard thermometer used should be graduated to tenths of a degree. Correct the reading by the following formula:

$$S = T + 0.000154n(T-t)$$

in which S = the corrected melting point, T = the observed melting point, n = the length of the mercury column above the surface of the liquid, expressed in degrees, and t = the temperature of the air about the mercury column as determined by a second thermometer.

Bömer states that by this method the analyst can detect in edible animal fats 1 to 2 per cent of oils rich in phytosterol (cottonseed, peanut, sesame, rape, hemp, poppy, and linseed), and 3 to 5 per cent of oils containing smaller amounts of this constituent (olive, palm, palm kernel, and probably cocoanut). He found the corrected melting point of cholesterol acetate to be 114.3° to 114.8° and of phytosterol acetate, 125.6° to 137.0° according to the source.

^{*} Zeits. Unters. Nahr. Genuss., 4. 1901, p. 1070.

CONSTANTS OF THE MOST COMMON EDIBLE FATS AND OILS AND THEIR ADULTERANTS.

Except when otherwise stated, the constants are mainly those of Lewkowitsch.

Oil or Pat.	Specific Gravity at 15.5° C.	Melting- point.	Solidifying- point.	Reichert- Meissl Number.	Saponifi- cation Number.	Jodine Number.	Bromine Index.	Insoluble Patty Acids, Hehner's Number.	Maumené Number.	Specific Tempora- ture Reac- tion.
Poppyseed oil	0.924 to 0.927		-16 to	0.0	190.1 to	133 to 143	0.835*	94.2 to 95.2	86 to 88	
Sunflower oil.	0.924 to		0 1	:	193.5	119 to		56	73	•
Corn oil.	0.9213 to			4 to \$	188 to	3	• • • • • • • • • • • • • • • • • • • •	93 to 96	81 to 86	6.381
Cottonseed oil	0.922 to	3 to 4		0.95	193 to	108 to	0.645	95 to 96	75 to 90	1.69 to
Sesame oil.	0.923 to			1.2	189 to	103 to	0.695	95.7	65.5	
Rape oil.	0.9237 0.9132 to		-2 to	. e.	170 to	94 to	0.632\$	95.1	55 to 64	1.25 to
Black mustard oil	0.9168 0.916 to		111	:	174	96 to	0.763\$	95.1	43	•
White mustard oil	0.920 0.914 to		- 8 to		170 to.	92 to		96.2	44 to 49	
Peanut oil	0.910 0.917 to		- 3 to o		190 to	83 to 97	0.530	8.8	45 to 51	tos to
Olive oil	0.9209 0.916 to		-6 to	9.0	185 to	79 to 103	o. 5 to	ᅿ	41.5 to 45	0.89 to
Lard oil	0.918 0.915 to		*		195 to	52.57 to	4	:	41 to 45**	*66
Cottonseed stearin.	0.9188 to	29 to 32	16 to 22		195	89 to		95.9	84	
Cocoa butter.	0.950 to	28 to 33	23 to 21.5	0.2 to 0.8	192 to	32 to		94.59		:
Cocoanut oil.	0.9115	21 to 24	22 to 14	7 to 8.4	246 to	8 to 9.5	:	88.6 to		:
Lard.	0.934 to	36 to 40.5	27.1 to	1.10	195.4	46 to 70	:	d di	24 to 27.5	3.71 to
Beef tallow.	0.043 to	40 to 45	35 to 27	0.5	192.2 to	35 to 47		93.6		
Mutton tallow	0.937 to	44 to 45	36 to 41	:	192 to	32 to 46	:	95.5	:	• • • • • • • • • • • • • • • • • • • •
Butter	0.926 to	88 to 33	20 to 23	25 to 30.4	222	26 to 38		86.5 to		
				7					7	

	11	Zeiss	1		Free Patty		Unsaponi-	Mix	Mixed Fatty Acids.	ds.
Oil or Fat.	Bromina- tion.	Refractom- eter Read- ing.	ture of Reading.	Acetyl Value.	Acids as Oleic.	Acid Value.	fiable Matter.	Solidifying- point.	Melting- point.	Iodine Number.
Poppyseed oil		63.4	004	13.1#	r fo to	0.7 to	0.43	16.2	20.5	130
		72.3	25°	:	27.71	11.0	0.31	:	22 to 24	121 to
Corn oil.	21.5	65.0	35°			1.35 to	1.7 to	:	18 to 20	119.5
Cottonseed oil	19.4	67.6 to	250	7.6 to		0.0	0.73 to	32 to 35	35 to 38	111 to
Sesame oil	23 to	6 89.4	250	11.5	0.4788 to	0.23	0.95 to	22.9 to	26 to 32	110.45
Rape oil.	23.9 17 to	89	25°	14.7	2.45	1.4 to	0.58 to	23.6 12 to 13	16 to 19	99 to 103
	30	59.5	°0 4			1.36 to			91	109.6
White mustard oil.		58.5	°¢			7.35 5.4			15 to 16	95.3
Peanut oil.		66 to	25°	3.41	1.45m to	1.2	0.54 to	29.3	27.7 to	96 to 103
Olive oil	1.5	62.4	250	10.64	8.287 I.55 to	1.9 to	0.46 to	17.2 to	24 to 27	86 to 90
Lard oil.		53.9	35°		MCC . 0	2		•		
Cottonseed stearin								35.1	27 to 30	\$
Cocoa butter		46 to	°	2.8		1.1 to		48.3	48 to 50	33 to 39
Cocognut oil.		34.8	° 0‡	0.9 to		1.85 5 to 50		22.5 to	25 to 27	8.4 to
Lard.	9 to	48.6 to	•	2.6	o.35 to	o. 54 to	0.23	41 to 42	43 to 44	•
Beef tallow	11.8 6.1 to	\$1.2	6 0 4	2.7 to	L100.1	1.28 3.5 to		37.9 to	43 to 44	41.3
Mutton tallow	7.2 7.6 to	\$	°0 1	ø :		1.7 to		40.15 to	49 to 50	34.8
Butter.	8.9 6.6 to 9.5	41 to	°0 	1.9 to 8.6	:	35.38		*	38 to 40	28 to 31
	* Levallois. † Gill and Hatch. ‡ Thompson and Balla. # Medicus and Scherer.	Levallois. Gill and Hatch. Thompson and Ballantyne. Medicus and Scherer.	me.	# Girarr # Hopk ## Gill. ## Vary.	Girard. ¶ Hopkins. ¶ Gill.	共三路管	## Benedict and Ulzer. Crossley and Le Suer. Shordlinger.	Ulzer. Le Suer.		

Numerous experiments, made both in Europe and America, show that feeding milch cows and swine with oil cakes does not introduce phytosterol into either the fat of the milk or the lard, although both fats may respond to the Halphen test, or give abnormally high Polenske numbers as a result of feeding with cottonseed or cocoanut cake respectively, and although the lard (not the butter fat) may respond to the Baudouin test, owing to feeding with sesame cake. (See pp. 531, 560).

Paraffin, sometimes present as an adulterant of fats, is best determined as follows: * Boil 2 grams of the fat with 10 cc. of 95% alcohol and 2 cc. of 1:1 sodium hydroxide solution, connect the flask with a reflux condenser, and heat for an hour on the water-bath, or until saponification is complete. Remove the condenser, and allow the flask to remain on the bath till the alcohol is evaporated off and a dry residue is left. Treat the residue with about 40 cc. of water and heat on the bath, with frequent shaking, till everything soluble is in solution. Wash into a separatory funnel, cool, and extract with four successive portions of petroleum ether, which are collected in a tared flask or capsule. Remove the petroleum ether by evaporation and dry in the oven to constant weight.

It should be noted that any phytosterol or cholesterol present in the fat would come down with the paraffin, but the amount would be so insignificant that with added paraffin actually present, it may be disregarded. The character of the final residue should, however, be confirmed by determining its melting-point and specific gravity, and by subjecting it to examination in the butyro-refractometer. The melting-point of paraffin is about 54.5° C.; its specific gravity at 15.5° is from 0.868 to 0.915, and on the butyro-refractometer the reading at 65° C. is from 11 to 14.5.

MICROSCOPICAL EXAMINATION OF OILS AND FATS.

Excepting in the case of solid fats, the use of the microscope has hitherto been comparatively restricted. In the examination of lard and butter for adulterants, the use of the microscope is often of great value, and will be described more fully under these special fats. In general the best fat crystals are obtained by slow crystallization at room temperature from an ether solution, or from a mixture of ether and alcohol. The first crystals formed may often with advantage be filtered out, and washed with the alcohol and ether mixture on the filter, dissolved finally in ether, and the latter allowed to evaporate spontaneously. The crystals are then examined in a medium of ether.

If it be desired to separate the liquid oleins from an oil, so that crystals of the solid fats are left for examination, Gladding* recommends dissolving the fat in a mixture of two volumes of absolute alcohol and one volume of ether in a test-tube, which is stoppered with cotton and set for half an hour in ice water, during which time the more solid stearin and palmitin will have crystallized out. This portion is then separated from the mother liquor by filtration through an alcohol-wet filter-paper, and the crystals finally treated as in the preceding section, being examined in a medium of olive or cottonseed oil.

OLIVE OIL.

Source.—Olive oil is derived from the fruit of the cultivated thornless olive tree, Olea Europea sativa,† of which there are a great many varieties, originally grown in Asia Minor, Greece, Palestine, and southern Europe, and now cultivated extensively in California, Peru, and Mexico, as well as in Australia. Most of the olive oil of commerce, especially of the choicest varieties, is supplied by southern France, Spain, and Italy. The tree is an evergreen of slow growth and great longevity.

The ripe olive fruit is purple or purplish black in color; it is round or oval in shape, and from 2.5 to 4 cm. in diameter. The oil is contained in the parenchyma cells of the fruit suspended in a watery fluid. A thick skin incloses the fruit, and within is a kernel, which itself contains oil. The fruit contains from 40 to 60 per cent of oil. According to Brannt, the average composition of the olive is as follows:

	Flesh,	Stone,	Seed,
	Per Cent.	Per Cent.	Per Cent.
Oil		5-75 85.89 2.50 4.16 4.20 100.00	12.26 79.38 2.16 6.20 100.00

Preparation.—The finest virgin oil is produced from hand-picked, peeled olives, from which the kernels or pits have been removed. A somewhat inferior grade of oil is produced from the whole olive including the pit, while a distinctly low grade oil is obtained from the stones, or kernels, which are ground into a coarse meal and subjected to pressure, or to the action of such solvents as carbon bisulphide.

^{*} Jour. Am. Chem. Soc., 1896, 18, p. 189.

[†] As distinguished for the wild thorny species, Europæa sylvestris.

¹ Animal abd Vegetable Fats and Oils.

In the process of manufacture the fruit, after first being dried, is reduced to a pulp in a stone or iron mill, and the pulpy mass, contained in baskets or bags, is subjected to pressure in an iron press. The very highest grade of virgin oil is that which runs out from the pulp with little or no pressure. After the first pressing, the pomace is ground, treated with water, and again subjected to pressure. Several pressings in this manner may be carried out, each yielding an oil inferior to that preceding, the lowest grades being used for lubricants and in the manufacture of soap.

Nature and Composition.—The better grades of olive oil, suitable for table and medicinal purposes, possess a pleasant, bland taste, and a distinctive and agreeable odor, unmistakable in character for that of any other oil. The finest virgin oil is pale green in color, due to the presence of chlorophyll, which is closely associated with the oil globules in the cellular tissue of the fruit. Some varieties of olive oil are nearly colorless, while others are a deep golden yellow.

Olive oil contains 28% of solid glycerides, chiefly palmitin and a very small amount of arachin, and 72% of liquid glycerides, mainly olein with a little linolein. Stearin is practically absent.

Lewkowitch* states that olive oil differs from most vegetable oils in containing cholesterol but not phytosterol.

Gill and Tufts † show, as a result of numerous experiments, that olive oil is not thus exceptional, but that the unsaponifiable alcohol is phytosterol and not cholesterol.

Olive oil is very soluble in chloroform, benzol, and carbon bisulphide, but is sparingly soluble in alcohol. Five parts of ether will dissolve 3 parts of the oil.

ADULTERANTS.—As a rule the low grade olive oils are most subject to adulteration, by reason of the fact that it hardly pays to destroy or even modify the fine quality and delicacy possessed by a first-class oil, which would inevitably be the result if even a small amount of foreign oil were added. Furthermore, if olive oil be slightly rancid or for any reason lacking in flavor, the admixture of a bland oil tends rather to minimize the fact.

The most common adulterant of olive oil in this country is naturally cottonseed oil, which is often substituted wholly for it. In Europe peanut oil is sometimes used both as an admixture and even as a substitute, since it possesses in itself a rather pleasant flavor, rendering it especially adapted for use as an adulterant. Other cheap oils used for this purpose are corn, mustard, poppyseed, rape, sesame, and sunflower

^{*} Chem. Anal. of Oils, Fats, and Waxes, 2d ed., p. 452.

[†] Jour. Am. Chem. Soc., XXV, 1903, p. 498.

oil. The writer has also found in samples of alleged olive oil sold in Massachusetts cocoanut oil* and even fish oil.

Pure Olive Oil of the U.S. Pharmacopæia.—The requirements of the Pharmacopæia are as follows:

Specific gravity, 0.910 to 0.915 at 25° C. (77° F.); iodine value not less than 80 nor more than 88; saponification value 191 to 195. Very sparingly soluble in alcohol, but readily soluble in ether, chloroform, or carbon disulphide.

When cooled to about 10° C. (50° F.), the oil should become somewhat cloudy from the separation of crystalline particles, and at 0° C. (32° F.) it should form a whitish, granular mass.

If 2 cc. of olive oil be shaken vigorously with an equal volume of nitric acid (sp. gr. 1.37), the oil should retain a light yellow color, not becoming orange or reddish brown, and after standing for six hours, should change into a yellowish-white solid mass and an almost colorless liquid (absence of appreciable quantities of cottonseed oil and other seed oils).

Olive oil should not show the cottonseed oil reaction with the Bechi and Halphen test, p.518, northe sesame oil reaction with the Baudouin test, p. 519.

U. S. Standards.—Olive oil is the oil obtained from the sound, mature fruit of the cultivated olive tree (Olea europoea L.) and subjected to the usual refining processes; is free from rancidity; has a refractive index (25° C.) not less than 1.4660 and not exceeding 1.4680; and an iodine number not less than 79 and not exceeding 90. Virgin olive oil is olive oil obtained from the first pressing of carefully selected, hand-picked olives

Reaction with Strong Acid.—Pure olive oil, when shaken or stirred with an equal volume of concentrated nitric or sulphuric acid, turns from a pale to a dark-green color in a few minutes. If, under this treatment, a reddish to an orange, or brown coloration is produced, the presence of a foreign vegetable oil (usually a seed oil) is to be suspected.

Bach gives the following table showing the action of strong nitric acid on various oils:

Kind of Oil.	After Agitation with Nitric Acid.	After Heating for Pive Minutes.	Consistency after Standing Twelve to Eighteen Hours.
Olive. Peanut. Rape Sesame Sunflower Cottonseed	" rose " " White Dirty white	Orange-yellow Brownish yellow Orange-yellow Brownish yellow Reddish yellow Reddish brown	Solid '' Liquid Buttery ''
Castor.	Pale rose	Golden yellow	

^{*} A sample of alleged olive oil purchased in a Massachusetts drug store and found to be adulterated with cocoanut oil, had the following constants:

Specific gravity 0.911 Iodine number 74.5 Reichert-Meissl number 2.90 Butro-refractometer at 26° 56.5 The Zeiss Butyro-refractometer furnishes one of the most useful and easily applied preliminary means of judging the purity of the sample. If the reading is beyond the limits of pure olive oil, it at once indicates adulteration and often points to the particular adulterant. On the other hand, it is not always safe to assume the oil to be pure if the reading is correct, since mixtures of higher and lower refracting foreign oils may be so skillfully prepared as to read well within the limits of the pure oil on the refractometer scale. The refractometer reading of pure cottonseed oil is almost five degrees higher than that of pure olive.

READINGS ON ZEISS REFRACTOMETER OF OLIVE AND COTTONSEED . OILS.*

Temperature (Centigrade).	Scale Reading.		Temperature	Scale Reading.	
	Olive Oil.	Cottonseed Oil.	(Centigrade).	Olive Oil.	Cottonseed Oil
35.0	57.0	61.8	25.5	62.4	67.5
34-5	57.2	62.1	25.0	63.0	67.9
34.0	57-4	62.3	24.5	63.3	68.2
33-5	57 - 7	62.5	24.0	63.6	68.5
33.0	58.0	62.8	23.5	63.9	68.8
32.5	58.3	63.0	23.0	64.2	69.1
32.0	58.5	63.2	22.5	64.5	69.4
31.5	59.0	63.6	22.0	64.8	69.7
31.0	. 59-2	64.0	21.5	65.1	70.0
30.5	59-4	64.2	21.0	65.4	70.3
30.0	59-9	64.5	20.5	65.7	70.6
29.5	60.1	64.9	20.0	66.o	70.9
29.0	60.3	65.1	19.5	66.3	71.2
28.5	60.6	65.3	19.0	66.6	71.5
28.0	60.9	65.7	18.5	66.9	71.8
27-5	61.1	66.0	18.0	67.2	72.1
27.0	61.5	66.5	17-5	67.5	72.4
26.5	62.0	67.0	17.0	67.8	72.7
26.0	62.2	67.3	16.5	68. z	73.0

The Elaīdin Test, in the case of pure olive oil, is very distinctive, since it yields by far the hardest elaïdin of all the common oils, and solidifies the most quickly.

Archbutt † shows the effect on this test of the mixture with olive oil of various proportions of rape and cottonseed oil, as follows:

Kind of Oil.	Minutes Required for Solid- ification at 25° C.	Consistency.
Olive oil	From 9 to 11½ hours	Hard but penetrable Buttery Very soft.

Ann. Rep. Mass. State Bd. of Health, 1899, p. 647. | Jour. Soc. Chem. Ind., 1897, p. 447.

Cottonseed Oil as an adulterant is best detected by means of the Halphen or Bechi tests. Its presence in notable quantities increases the specific gravity, refractometer reading, and iodine number very materially. Its high Maumené figure is also distinctive.

Peanut Oil, when present to a considerable extent, betrays its presence by its peculiar bean-like flavor. Most of the constants of peanut oil lie within the limits of olive oil, with the exception of the higher rodine number and refractometer reading. A considerable admixture of peanut oil raises the refractometer reading perceptibly over that of pure olive. Its presence is best shown positively by tests for arachidic acid (p. 523), noting that traces of arachin have been reported in pure olive oil, insufficient, however, to interfere with the detection of added peanut oil.

Sesame Oil differs more particularly from olive in its higher specific gravity and iodine and Maumené numbers, and is readily detected by distinctive color tests (p. 510).

Rape Oil is characterized by a much lower saponification value and higher iodine number than olive.

Corn Oil differs materially from olive in its exceedingly high iodine number and refractometer reading. Its specific gravity and saponification numbers are also higher.

Lard Oil, when present in considerable quantity, is often rendered apparent by its characteristic odor on warming. Its low refractometer reading and iodine number are also distinctive.

Poppyseed Oil differs most widely from olive oil in its refractometric reading, its high dispersion, and its Maumené number, which in the case of poppyseed is 87° and of olive about 42°.

Cocoanut Oil in mixture with olive perceptibly raises the solidifying-point. When more than 12% of cocoanut oil is present, the sample will become solid when placed in ice water.

Fish Oils, when present, are rendered apparent by reason of their strong taste and smell, and by their very high iodine number. Boiling the sample with sodium hydroxide develops a peculiar reddish coloration, when fish oils are present.

Routine Examination of Olive Oil for Adulterants.—First note the smell and taste of the sample, and then take the refractometric reading. An abnormally high refraction indicates adulteration. Then test with strong nitric acid (p. 513). If the refraction is normal, and the color resulting from the acid reaction a pale green, the presumption is that the oil is pure. Test first for cottonseed oil by the Halphen reaction,

and then in succession try the various color reactions for sesame and rape oils. If all these are absent, and, by abnormal constants, or by color with nitric acid, there is reason to believe the oil is adulterated, determine carefully such of the constants as are most indicative, by their wide variation from olive, of poppyseed, mustard, and corn oils.

If all these oils are presumably absent, and either a high refractometer reading or a color reaction with nitric acid still indicates adulteration, peanut oil is more than likely to be present, and should be tested for either by Renard's or Bellier's method.

The edible oils and adulterants are arranged in order of their relative price about as follows:

Olive oil.

Peanut oil.

Lard oil.

Sesame oil.

Poppyseed oil.

Rape oil.

Corn oil.

Cottonseed oil.

COTTONSEED OIL.

Source and Preparation.—This oil, largely used as a table oil and as an adulterant of olive oil, is derived from seeds of the various species of the cotton plant, Gossipium, of which the most common are G. herbaceum, native to Asia, but cultivated extensively in southern Europe and in the United States, G. arboreum, in Asia and Africa, and G. barbadense, in the West Indies. G. religiosum and hirsutum are varieties of G. herbaceum.

The seeds are in reality a by-product in cotton manufacture. In shape they are irregularly oval, measuring from 5 to 8 mm. greatest diameter. The seed skin or pod is covered with the fiber of the cotton.

The seeds are first cleaned and separated from dirt by sifting machines, and from the fiber by specially constructed gins, after which they are cut into small pieces, freed from their hulls, crushed between rollers, and afterward submitted to hydraulic pressure in bags to express the oil, which is clarified by filtration or refined. The refining consists in washing the crude oil with sodium hydroxide solution, whereby the impurities are dissolved and thus removed.

Nature and Composition of Seeds and Oil.—The seeds of the cotton plant are rich in oil, containing from 10 to 29 per cent, according to the

variety.	Four	samples	of	American	cottonseed	were	found	to	be	com-
posed as	follow	s, accord	ding	g to Brann	ıt:*					

Constituents.	South Carolina.	Georgia I.	Georgia II.	Georgia III.
Water Cottonseed oil Nitrogenous compounds. Ammonia-making compounds. Gum, sugar, and soluble starch Cellulose, starch, and resin. Ligneous tissue	9.5 20.1 17.8 2.3 .8 26.2 17.6	10.1 16.2 17.4 2.9 .9 27.4	9.8 17.1 17.2 3.2 .7 26.1	8.2 19.6 18.1 3.7 .9 20.7 22.4
Ash (phosphate of lime, silica, alumina, iron, magnesia, potash, soda, etc.)	5-7	5-9	6.1	· 6.4
İ	100.0	100.0	100.0	100.0

Refined cottonseed oil is a pale-yellow oil of thick consistency, possessing a bland though pleasant taste and odor. It consists of the glycerides of oleic, stearic, palmitic, and linoleic acids, and evidently also a small content of hydroxyacids, though this has not been investigated as yet.

On cooling the oil to a temperature below 12° C. particles of solid fat will separate. At about 0° to -5° C. the oil solidifies. When the oil is brought in contact with concentrated sulphuric acid, a dark, reddish-brown color is instantly produced.

U. S. Standards.—Cottonseed oil is the oil obtained from the seeds of cotton plants and subjected to the usual refining processes; is free from rancidity, has a refractive index (25° C.) not less than 1.4700 and not exceeding 1.4725; and an iodine number not less than 104 and not exceeding 110.

"Winter-yellow" cottonseed oil is expressed cottonseed oil from which a portion of the stearin has been separated by chilling and pressure, and has an iodine number not less than 110 and not exceeding 116.

Cottonseed Stearin.—This product, used as an adulterant of lard as well as a substitute therefor, is obtained as a by-product in the manufacture of winter-yellow cottonseed oil. It is a light yellow fat, resembling butter in consistency.

Bechi's Silver Nitrate Test.—Hehner's Modification.—Two grams of silver nitrate are dissolved in 200 cc. of 95% alcohol free from aldehyde, 40 cc. of ether are added, and the reagent made very slightly acid with nitric acid.

In applying the test, a small quantity of the melted fat or oil is mixed in a test-tube with half its volume of the above reagent, and the tube is immersed in boiling water for fifteen minutes. With proper precautions

^{*} Vegetable Fats and Oils, p. 223.

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the presence of cottonseed oil is indicated by a more or less strong reduction of the silver, while an oil or fat free from cottonseed oil causes no appreciable reduction.

Certain oils free from cottonseed that have become rancid or decomposed, as well as fats that have been subjected to a high temperature, sometimes show a slight reduction with Bechi's test. In cases of doubt it is well to apply the test on the fatty acids as follows:

Milliau's Modification of Bechi's Test.*—Heat 20 grams of the sample with 30 cc. of alcoholic potash solution (20% potassium hydroxide in 70% alcohol), shaking at intervals till saponification is complete. Continue the heating for some minutes afterward until the alcohol is driven off, and dissolve the soap in 250 cc. of hot water. Add a slight excess of 10% sulphuric acid, and wash the separated fatty acids three times by decantation with water. Then proceed with a portion of the fatty acids as in Bechi's test.

Halphen's Test.—This is a much more delicate test for cottonseed oil than either of the preceding, as little as 2% of cottonseed oil being rendered apparent in olive oil. A mixture is made of equal volumes of amyl alcohol and carbon bisulphide in which 1% of sulphur has been dissolved. From 3 to 5 cc. of melted fat are mixed with an equal volume of the above reagent in a test-tube, loosely stoppered with cotton, and heated in a bath of boiling saturated brine for fifteen minutes. If cottonseed oil is present, a deep-red or orange color is produced. In its absence little or no color is developed.

Previous heating of the oil diminishes the delicacy of the Halphen test, and Holde and Pelgry † state that if cottonseed oil has been heated at 250° for ten minutes, it will fail to respond to the test. Fulmer finds that it is necessary to heat to 265 to 270° to render it wholly inactive to the test.

SESAME OIL.

Sesame or benné oil is pressed from the seeds of Sesamum indicum and S. orientale, both of which are now regarded as varieties of the same species, and S. radiatum. These plants are native to southern Asia, but now cultivated in nearly all tropical countries. The larger portion of commercial sesame oil is manufactured in England, France, Germany, and Austria.

The seeds are yellow to dark brown, and in some cases black, inclined to the oval in form, the average longest diameter being about 4 mm.

^{*} Monietur Scientifique, 1888, p. 366. † Jour. Soc. Chem. Ind., 18, 1899, p. 711.

The seeds are commonly subjected to cold pressure once, and afterwards twice pressed when warm, thus yielding three grades of oil. From 47 to 60 per cent of oil is contained in the seeds.

According to	Brannt* the	composition of	of sesame	seeds is	s as	follows:
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		mum ntale.		mum cum.
Oil. Organic substances. Protein therein. Nitrogen therein. Ash Water	55.63 30.95 7.52 3.90 100.00	21.42 3·39	50.84 35-25 6.85 7.06 100.00	22.30 3.56

Sesame oil consists of the glycerides of oleic, stearic, palmitic, and myristic acids. It is golden yellow in color, free from odor, and possesses a delicate and characteristic flavor, on account of which the highest grades are by some considered equal to olive oil as a condiment. It is accordingly sold to some extent as an edible oil. It was formerly used as an adulterant of olive oil, but has of late years been largely displaced by cheaper oils for purposes of adulteration. When cooled to -3° C., sesame oil congeals to a yellowish-white mass. Concentrated sulphuric acid converts it into a brownish-red jelly.

U. S. Standards.—Refractive index (25°) 1.4704 to 1.4717; iodine number 103 to 112.

Adulterants to be looked for in sesame oil are cottonseed, poppy-seed, corn, and rape oils.

Tocher's Test.†—One gram of pyrogallic acid is dissolved in 15 cc. of concentrated hydrochloric acid and mixed with 15 cc. of the sample in a separatory funnel. After standing for a minute, the aqueous solution is withdrawn and boiled. If sesame oil is present, the solution shows a red coloration by transmitted, and blue by reflected, light.

Baudouin's Test.‡—Dissolve 0.1 gram of cane sugar in 10 cc. of hydrochloric acid (specific gravity 1.20) in a test-tube, and shake thoroughly with 20 grams of the oil to be tested for one minute. Then allow the mixture to stand. The aqueous solution quickly separates from the oil, and in the presence of 1% or more of sesame oil will be colored deep red.

Certain pure Tunisian and Algerian olive oils have been found to cause a slight coloration with this test, but of a different shade from sesame. Moreover, if the test is applied to the fatty acids, no coloration in the case of olive oil is produced, while with sesame the color is the same as with the oil-

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^{*} Vegetable Fats and Oils, p. 251. † Chem. Zeit. Rep., 5, 1891, p. 15. ‡ Zeits. angew. Chem., 1892, p. 509.

Villivecchia and Fabris Test.*—This test was suggested on account of the fact that the color reaction in the Baudouin test was attributed to the agency of the levulose produced by the inversion of the sugar by hydrochloric acid. As furfurol is the chief product of the reaction between levulose and hydrochloric acid, it was substituted as follows: Dissolve 2 grams of furfurol in 100 cc. of 95% alcohol, and shake 0.1 cc. of this solution in a test-tube with 10 cc. of the oil to be tested and 10 cc. of hydrochloric acid (specific gravity 1.20) for half a minute. The aqueous layer, on settling out, will be colored deep red, if sesame is present.

Or 0.1 cc. of the alcohol furfurol solution is mixed with 10 cc. of oil and 1 cc. of hydrochloric acid in a separatory funnel, shaken well, and the separation aided by the addition of chloroform, which causes the aqueous layer, showing color with sesame oil, to float.

Since furfurol produces with hydrochloric acid alone a violet coloration, it is necessary to use it in dilute solution as above.

RAPE OIL.

Rape or colza oil is expressed from the seeds of the Brassica or rapeplant, of which there are three principal varities, Brassica napus, B. campestris, and B. rapa, one or another of which are cultivated in nearly every country of Europe, excepting Greece. Large amounts are also grown in India and China. The seeds are small, round grains, from 2 to 2.5 mm. in diameter, yielding from 30 to 45 per cent of oil. The seeds, according to Brannt, have the following average composition:

	Fresh Seeds.	Old Seeds.
Oil	36.80 49.30	38.50 53.25
Nitrogen therein	2.50 4.80	3.90
Water.	9.10	4.35
	100.00	100.00

In the process of preparation the seeds are first crushed, and the oil removed by pressing or extraction. The crude oil is of a brownish-yellow color, and when fresh is almost free from taste and smell, so that it serves, when cold pressed, as an edible oil, or an adulterant of such oils. It develops a disagreeable and peculiar taste and odor on long standing, due to the presence of certain albuminous and mucilaginous substances which it contains. These may be removed by refining, usually by treatment with sulphuric acid, but the refined oil has an unpleasant taste and odor.

^{*} Jour. Soc. Chem. Ind., 1894, pp. 13-69.

[†] Vegetable Fats and Oils, p. 240.

The principal components of rape oil are the glycerides of stearic, oleic, erucic, and rapic acids. The chief adulterants are cottonseed and poppyseed oils.

Palas Test for Rapeseed Oil.*—Mix in the cold 30 cc. of a 1% solution of fuchsin, 20 cc. of sodium bisulphite (specific gravity 1.31), 200 cc. of water, and 5 cc. sulphuric acid. If the sample of oil to be tested be shaken with the reagent, a rose-red coloration is obtained in the presence of rape oil, said to be delicate to the extent of detecting 2% of the oil in mixtures.

CORN OR MAIZE OIL

Corn oil is derived from the seed of the American grain Zea mays, or Indian corn, the constitution of the yellow and white varieties of which is, according to Andés,† as follows:

	Yellow	Corn,	White Corn,		
	Per C	ent.	Per Cent.		
Organic matter Starch Albuminoids. Ash Water Oil.	82.93 1.32 9.50 6.25	61.95 10.71	80.76 1.04 10.60 7.60 100.00	62.23 9.62	

Nearly all the oil is contained in the germ of the seed, the oil constituting in fact over 20% of the germ. Corn oil consists chiefly of the glycerides of palmitic and oleic acids. There is some doubt as to the presence of stearin. It is golden yellow in color, and possesses a pleasant odor and taste, resembling in flavor freshly ground grain.

It is prepared by subjecting to hydraulic pressure the germ separated in the manufacture of starch and of glucose, the germs yielding about 15% of pure oil. While most of the oil of commerce is a by-product from starch and glucose factories, a small amount is recovered from the residue of fermentation vats in the manufacture of alcohol. Corn oil is coming to be used more and more as an adulterant of olive oil, and, according to Lewkowitsch, of lard.

It is claimed by Hopkins,‡ by Hoppe-Seyler, and others, that corn oil,

^{*} Analyst, XXII, p. 45.

[†] Vegetable Fats and Oils, p. 131.

¹ Jour. Am. Chem. Soc., 1898, 20, p. 948.

unlike most vegetable oils, contains cholesterol. Olive oil was long supposed to be unique as a vegetable oil in containing this substance. Hopkins, on the assumption that cholesterol occurs in corn oil, suggested that a test for corn oil as an adulterant of certain vegetable oils lay in the identification of cholesterol.

Gill and Tufts * claim that, while the alcohol of corn oil is not phytosterol, neither is it cholesterol, but a third substance, known as sitosterol,† occurring in wheat and rye.

There are no color reactions identifying corn oil as such. Its presence in other oils is indicated only by its influence on the various constants, the iodine number and refractometric reading especially being much higher than those of other edible oils.

PEANUT OIL.

Peanut or arachis oil is obtained from the seeds of the Arachis hypogaa (peanut, ground nut, or earth nut) cultivated in most tropical countries, notably in South America, China, India, and Japan. The plant is a creeping herb, developing its blossoms in the axes of the leaves. The fruit buds grow down into the earth, where the fruit is ripened, forming the well-known peanuts of commerce, the composition of which, according to Brannt, is as follows:

Oil	Per Cent. 37-48 52.86 27.25 2.43 7-37	" 2-50 " 2-75
	100.00	100.00

Peanut oil is composed chiefly of the glycerides of oleic, palmitic, hypogæic, and arachidic acids. The oil is extracted by pressure, the first cold-drawn oil being practically colorless, and possessing a pleasant taste suggestive of kidney beans. It is especially adapted for use as a salad or table oil. A second pressure of the moistened residue from the first yields an inferior oil, yellowish in color, also somewhat used for edible purposes, and sometimes commercially called "butterine oil."

U. S. Standards.—Refractive index (25°) 1.4690 to 1.4707; iodine number 87 to 100.

^{*} Jour. Am. Chem. Soc., XXV, 1903. † Burian, Monatsh. Chem., 18, 1897, p. 551.

Adulterants of peanut oil are cottonseed, poppyseed, rape, and sesame oils. Very little pure peanut oil is found in commerce in the United States. It is to be looked for as an adulterant of French and Italian olive oils.

Characteristic Tests.—Peanut oil, when pure or nearly pure, may as a rule be readily identified from other oils. When present in large admixture in other oils it is not difficult to detect, but when only a small amount is present, in olive oil for instance, its detection becomes a more troublesome matter.

This difficulty arises from the fact that the constants of peanut oil are nearly the same as those of olive, with the single exception of the refractometric reading. Furthermore, there is no readily applied color test identifying peanut oil.

All the other common adulterants of olive oil, as cottonseed, sesame, corn, poppyseed, and rape oils, are readily identified, when present in small amounts, either by special color tests, or by reason of the fact that certain of their constants differ very widely from those of olive oil. Much more care and precaution are necessary in dealing with small admixtures of peanut oil than with almost any other adulterant.

The Renard Test* has long been in use for detecting and estimating peanut oil in mixtures. In its original form this test did not give entirely satisfactory results, and earlier led to some erroneous conclusions. In recent years, however, it has been so modified and improved as to be capable of quite positive results when carefully carried out. While arachin is said to occur in minute traces in olive oil, its presence is not sufficiently marked to interfere with the use of the Renard method in detecting any decided admixture of peanut oil.

The following modification of the Renard method, devised by Tolman,† has been adopted by the A. O. A. C.:

Twenty grams of the oil are saponified in a 250-cc. Erlenmeyer flask with 200 cc. of alcoholic potassium hydroxide (40 grams potassium hydroxide in 1 liter of 95% redistilled alcohol). Neutralize with dilute acetic acid, using phenolphthalein as an indicator, and wash into a 500-cc. flask containing a boiling mixture of 100 cc. water and 120 cc. 20% solution of lead acetate.

Boil for a minute and cool the contents of the flask by immersing in cold, or, preferably, ice water, whirling the flask occasionally so that

^{*} Comp. Rend., 73, 1871, p. 1330.

[†] U. S. Dept. of Agric., Bur. of Chem., Bul. 65; also Bul. 77, and Bul. 107 (rev.).

the soap when cold adheres to the sides of the flask. The water and excess of lead acetate can then be poured out, leaving the soap in the flask. Wash by shaking and decantation, first with cold water and then with 90% alcohol. Add 200 cc. of ether, cork the flask, and allow to stand with occasional shaking till the soap is disintegrated, after which boil on a water-bath under a reflux condenser for five minutes. Cool the soap solution down to a temperature between 15° and 17°, and allow it to stand for about twelve hours.

Filter and thoroughly wash the precipitate with ether, after which the soap in the filter is washed back into the original flask with a stream of hot water acidulated with hydrochloric acid.

Add an excess of dilute hydrochloric acid, partially fill the flask with hot water, and heat until fatty acids form a clear oily layer. Fill the flask with hot water, allow the fatty acids to harden and separate from the precipitated lead chloride, wash, drain, repeat washing with hot water, and dissolve the fatty acids in 100 cc. of boiling 90% by volume alcohol. Cool to 15° C., shaking thoroughly to aid crystallization.

From 5 to 10 per cent of peanut oil can be detected by this method, as it effects a complete separation of the soluble acids from the insoluble, which interfere with the crystallization of the arachidic acid. Filter, wash the precipitate twice with 10 cc. of 90% alcohol, and then with 70% alcohol. Finally dissolve off the precipitate with boiling absolute alcohol, evaporate to dryness in a tared dish, dry and weigh. To the weight add 0.0025 gram for each 10 cc. of 90% alcohol used in the crystallization and washing, if done at 15° C., and 0.0045 gram for each 10 cc. if done at 20°. The approximate amount of peanut oil is found by multiplying the weight of arachidic acid by 20.

Arachidic acid crystals thus obtained should be examined microscopically. The melting-point should lie between 71° and 72° C.

Methods of J. Bellier.*—Qualitative Test.—Saponify I gram of the oil with 5 cc. of an alcoholic potash solution containing 85 grams potassium hydroxide per liter of strong alcohol, conducting the saponification in a small Erlenmeyer flask on the water-bath. After saponification, boil for two minutes, neutralize with dilute acetic acid, using phenolphthalein as an indicator, and cool by setting the flask in water at a temperature of from 17° to 19°. After a short time, a precipitate nearly always comes down. Then add to the solution 50 cc. of 70% alcohol, containing 1% by volume of strong hydrochloric acid (specific

^{*} Ann. Chim. Anal., 1899, 4, p. 49; Zeits. für untersuch. Nahr., 1899, 2, p. 726.

gravity 1.20). Cork the flask, shake vigorously, and again cool by setting the flask in the above cooling-bath. In the absence of a precipitate, the oil may be pronounced free from peanut. If 10% or more of peanut oil is present, a more or less characteristic precipitate forms, and often with less than 10% a cloudiness in the solution is perceptible after standing between 17° and 19° for half an hour. Pure olive oil remains perfectly clear as a rule.

A few varieties of olive oil from Tunis especially high in solid fat acids, as well as cottonseed oil and sesame oil, give similar turbidity on the addition of the 70% alcohol. To distinguish between these oils and peanut oil, heat the mixture on the water-bath till complete solution takes place, and again cool to 17° to 19° . In the case of peanut oil the cloudiness or precipitate again occurs to the same extent as before, while in the other cases the solution should remain clear or nearly so.

Quantitative Determination.—Saponify 5 grams of the oil with 25 cc. of the above alcoholic potash solution in a 250-cc. Erlenmeyer flask, neutralize exactly with acetic acid, and cool quickly in water. After standing an hour, pour upon a 9-cc. filter and wash the precipitate with 70% alcohol containing 18% by volume of hydrochloric acid, the temperature of the solution being not less than 16° nor more than 20°. Continue the washing till the wash water no longer shows turbidity when diluted with water.

Dissolve the precipitate in 25 to 30 cc. of hot 95% alcohol, dilute with water until the alcohol is 70%, let stand in water at 20°, filter, wash with 70% alcohol, dry at 100°, and weigh.

Bellier states that he has recognized with certainty as small an admixture as 2% of peanut oil by this method.

MUSTARD OIL.

The fixed oil of mustard is a by-product expressed from the seeds of the black and white mustard (Sinapis nigra and S. alba) in the process of preparation of mustard flour as a spice. The seeds contain from 25 to 35 per cent of oil.

Mustard oil somewhat resembles rape in composition, containing glycerides of erucic, behenic, and probably rapic acid.

Black mustard oil is brownish yellow in color, having a mild flavor, and an odor but slightly suggestive of mustard. White mustard oil is golden yellow, and has a somewhat sharp taste.

Mustard oil is an alleged adulterant of edible oils, though by no means a common one.

POPPYSEED OIL.

This oil is obtained from the seeds of the opium poppy (Papaver somnijerum), native in the countries east of the Mediterranean, and cultivated extensively for opium and for oil in all parts of Europe, Asiatic Turkey, Persia, Egypt, India, and China. Most of the oil of commerce comes from France and Germany.

There are two chief varieties of poppy, the black (P. nigrum) and the white (P. album), the finest oil being produced from the white. The seeds are somewhat flattened in form and kidney-shaped, yielding from 40 to 60 per cent of oil. According to Brannt the seeds have the following composition:

	White Poppy- seed.	Black Poppy- seed.		
Oil. Organic substances Protein therein Ash Water	55.62 32.11 16 89 3.42 8.85	51-36 35-14 17-50 4.00 9-50		

The oil is obtained by crushing the seeds and applying pressure. The best grade of cold-drawn oil is pale yellow in color, possessing a pleasant taste when fresh, and being practically free from odor. Lower grades shade into deeper yellow and even reddish color, possessing a strong taste and odor. Poppyseed oil is much used in Europe as a table oil, and does not readily turn rancid. It is composed of the glycerides of stearic, palmitic, and linoleic acids. Poppyseed oil has been used to some extent as an adulterant of olive oil. It is itself not infrequently adulterated with sesame oil.

SUNFLOWER OIL.

Sunflower oil is derived from the seed kernels of the plant of the same name (*Helianthus annuus*), originally grown in Mexico, but now cultivated most extensively on a commercial scale in southern Russia.

	According	to	S.	M.	Babcock *	the	composition	of	sunflower seeds
is	as follows:								

	Air-dry.	Dried.
Water	12.68	
Ash	3.00	3.43
Albuminoids (N×6.25)	15.88	3.43 18.19
Crude fiber	29.21	33-45
Nitrogen-free extract	18.71	21.43
Fat (ether extract)	20.52	23.50
	100.00	100.00

The seeds are long, black, and oval in shape, yielding from 18 to 28 per cent of oil. The liquid fatty acids of sunflower oil consist for the most part of linoleic, but little oleic acid being found.

The seeds are first shelled, then crushed, and finally submitted to pressure both cold and hot.

Sunflower oil is pale yellow in color, has a mild, pleasant taste, and is nearly free from odor. The cold-drawn oil is the variety most used for edible and culinary purposes in Russia, and as an adulterant of olive oil. Its use as an adulterant is, however, limited, and the writer has no knowledge of its having been found in olive oils used in the United States.

ROSIN OIL.

Rosin oil is prepared by the distillation of common rosin, and is an alleged adulterant of olive oil. It may be detected when present by shaking 1 to 2 cc. of the sample with acetic anhydride while warming. Cool, remove the anhydride by a pipette, and add a drop of sulphuric acid (specific gravity 1.53). Rosin oil gives a fugitive-violet color.†

Cholesterol also responds to this color reaction.

Renard's Test for Rosin Oil.—Prepare a solution of stannic bromide by allowing dry bromine to fall drop by drop upon tin in a dry, cool flask, and dissolving the product in carbon bisulphide.

Add a drop of this reagent to 1 cc. of the oil. In presence of rosin oil a violet color will be produced.

Polarization Test for Rosin Oil.†—The oil is dissolved in definite proportion in petroleum ether, and polarized in a 200-mm. tube. Rosin

^{*} The Sunflower Plant, its Cultivation, Composition, and Uses. U. S. Dept. of Agric., Div. of Chem., Bul. 60, p. 18.

[†] U. S. Dept. of Agric., Bur. of Chem., Bul. 65, p. 32.

oil polarizes from +30 to +40 on the cane sugar scale, while other oils have a reading between +1 and -1.

COCOANUT OIL.

Cocoanut oil is the fat expressed from the kernels of the cocoanut or fruit of the cocoa palm (*Cocos nucifera*), indigenous to the South Sea Islands and to the East-Indian archipelago, but grown in many tropical countries.

It is sometimes known as "copra oil," from the copra or pulp, which contains from 60 to 70 per cent. of fat. According to Brannt, the composition of the pulp is as follows:

	Indian Copra.	African Copra.		
Oil,	68.75	66.80		
Organic substances	23.65	25.25		
Albuminous substances	9.16	10.20		
Ash	1.45	1.50		
Water	6.15	1.50 6.45		
				
	100.00	100.00		

In the preparation of the oil the moist copra is separated from the shell, crushed in mortars and subjected to pressure, yielding a milky mass. This is then heated in boilers, and the oil which rises to the surface is removed by skimming.

In some localities the pulp is first dried and then pressed.

Cocoanut oil is usually white and possesses a mild taste and pleasant odor. The cold-drawn Malabar oil is of greenish color, and is used by the natives as an edible oil or substitute for butter. This variety is seldom found in commerce.

Cocoanut oil contains, besides palmitin and olein, large proportions of myristin and laurin. Unlike the other vegetable oils, it contains also notable quantities of the glycerides of the volatile fatty acids caproic, capric, and caprylic, hence the high saponification value and Reichert number. The most characteristic constant is the Polenske number. The oil is rarely adulterated.

Cocoanut oil easily becomes rancid. According to Andés, crystals of cocoanut oil appear under the microscope as a thick network of long needles.

COCOA (CACAO) BUTTER.

This preparation is not, properly speaking, in itself an edible fat. It is a by-product in the manufacture of cocoa, being removed by pressure from the crushed and ground cocoa nibs. The fat in cocoa beans varies from 36 to 50 per cent. The expressed fat is yellowish white, of a tallow-like consistency, has a pleasant taste and an odor suggestive of chocolate. It keeps a long time without turning rancid. In composition it consists of the glycerides of stearic, palmitic, and lauric acids, with traces of the glycerides of arachidic and butyric acids.

Its demand for pharmaceutical purposes is, however, sufficiently great to render the use of cocoa-butter as an adulterant of food-fats extremely rare. It should be borne in mind as a possible adulterant in examining various oils.

It is subject to adulteration with paraffin, tallow, and cottonseed stearin.

TALLOW.

The rendered fats of various animals, especially the cow and sheep, constitute what is generally known as tallow. The untreated fatty tissues are more properly known as suet, the tallow being the clear fat separated entirely by heat from the cellular material.

Tallow consists almost entirely of olein, palmitin, and stearin. Mutton tallow is usually, but not always, harder than beef tallow.

Excepting in the manufacture of material for oleomargarine, wherein the heart and caul fats of beef are almost exclusively used, the fats from different parts of the animal are not, as a rule, separated.

Fresh tallow has very little free fatty acid, but when it becomes rancid, the fat contains sometimes as high as 12% of free acid, reckoned as oleic.

Tallow is of chief interest to the food analyst in connection with its use as an adulterant of lard.

BUTTER.

Nature and Composition.—Butter is the product obtained by the churning of cream or milk, whereby the fat particles are caused to adhere together into a compact mass, inclosing a certain portion of the casein, the excess of milk serum being subsequently largely removed by washing and mechanical working.

Butter fat is of extremely complex composition, containing a larger variety of glycerides than any other fat. Besides olein, palmitin, and stearin, the usual glycerides of the insoluble or fixed fatty acids found in most fats, butter contains notable quantities of the glycerides of a number of the volatile fatty acids, chief among which are butyrin, caproin, caprin, and caprylin, to which are due its distinctive taste, and which by exposure to light and air readily become decomposed into their fatty acids—butyric, caproic, capric, and caprylic, respectively. This decomposition in butter causes, or, more properly speaking, accompanies, what is commonly known as "rancidity."

The process of separation of butter fat into its component glycerides is a matter of extreme difficulty, and results obtained by different chemists vary widely. Separation has been attempted by fractional distillation, by methods depending on the difference in chemical affinity of the various acids, and on the difference in solubility of the various lower homologues in water at different temperatures.*

According to Browne, the composition of butter fat is as follows:

Acid.	Percentage of Acid.	Percentage of Triglycerides.
Dioxystearic	1.00	1.04
Oleic	32.50	33-95
Stearic	1.83	1.91
Palmitic	38.61	40.51
Myristic	g.8g	10.44
Lauric.	2.57	2.73
Capric	0.32	0.34
Caprylic	0.49	0.53
Caproic		2.32
Butyric	5-45	6.23
-		
Totals	94-75	100.00

Upwards of 300 analyses of butter are summarized by König in the following table:

	Water,	Fat,	Casein,	Milk,	Sugar,	Lactic Acid,	Salts,
	Per Cent.	Per Cent.	Per Cent.	Per Cent.	Per Cent.	Per Cent.	Per Cent.
Minimum Maximum Mean		69.96 86.15 84.39	0.19 4.78 0.74	0.50	0.45 1.16	0.12	0.02 15.08 0.66

^{*} Browne, A Contribution to the Chemistry of Butter Fat, Jour. Am. Chem. Soc., 21, 1899, p. 807.

Effects of Feeding Oil Cakes on the Composition of Butter.—Experiments have shown that the substance which causes cottonseed oil to respond to the Halphen test passes into the milk fat on feeding cows with cottonseed cake, but the substance that gives the Baudouin reaction is never carried into the milk on feeding with sesame cake. A number of investigators have found that feeding with cocoanut cake raises somewhat the Polenske number of the milk fat. There is good evidence, however, that, while the addition of vegetable oils to butter introduces phytosterol, as detected by Bömer's phytosterol acetate test, this substance can not be introduced into the milk fat by feeding. These facts should be borne in mind in the examination of butter for foreign fats.

ANALYSIS OF BUTTER.

Preparation of the Sample.—A. O. A. C. Method.*—If large quantities of butter are to be sampled, a butter-trier or sampler may be used. The portions thus drawn, about 500 grams, are to be perfectly melted in a closed vessel at as low a temperature as possible, and when melted, the whole is to be shaken violently for some minutes till the mass is homogeneous, and sufficiently solidified to prevent the separation of the water and fat. A portion is then poured into the vessel from which it is to be weighed for analysis, and should nearly or quite fill it. This sample should be kept in a cold place till analyzed.

Water.—A. O. A. C. Method.—About two grams of the sample are weighed in a flat-bottomed platinum dish, such as is used for determining water in milk, and the dish and its contents kept in contact with the live stream of a water-bath till a constant weight is attained.

Patrick's Rapid Method.†—This method is especially suited for the use of dairymen, inspectors and others not provided with laboratory facilities.

Ten grams of the thoroughly mixed butter are weighed into a 250-cc. aluminium beaker, which, together with a glass rod has been previously tared, and boiled over (but not in) the flame of an alcohol lamp provided with a conical asbestos chimney, holding the beaker by means of a wire clamp in a nearly horizontal position to avoid loss from spattering or foaming, and whirling constantly to prevent overheating. The rod serves to break up lumps of curd which form, thus facilitating the drying.

^{*} U. S. Dept. of Agric., Bur. of Chem., Bul. 46, p. 43; Bul. 107 (rev.), p. 123.

[†] Jour. Am. Chem. Soc., 28, 1906, p. 1611; 29, 1907, p. 1126.

The heating should be so conducted as to avoid any considerable dis-

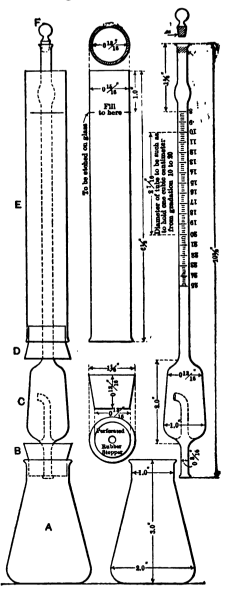


Fig. 100.—Gray's Apparatus for the Rapid Determination of Water in Butter.

as to avoid any considerable discoloration of the curd. With suitable heating the water may be removed in less than 15 minutes, after which the beaker is cooled in water and weighed. A balance sensitive to 10 milligrams, such as is used in weighing cream for testing by the Babcock method, is sufficiently accurate for weighing the butter.

Gray's Method.*—1. The Special Apparatus for this method, shown in Fig. 100, consists of a flask (A) connected by a close-fitting rubber stopper (B) with a graduated tube (C), and this in turn with a condenser jacket (E) by a rubber stopper (D). The tube C is closed by a glass stopper, the zero mark being the end of the stopper. Each mark of the graduation represents 0.02 cc. or, when 10 grams of butter are used, 0.2%.

2. Process.—Weigh 10 grams of the well mixed butter on a piece of parchment paper 13 cm. square, introduce into the flask, and add 6 cc. of a mixture of 5 parts of amyl acetate and 1 part of amyl valerianate, free from water-soluble impurities. Connect the apparatus as shown in Fig. 100, fill the condenser jacket with cool water to within 2.5 cm. of the top, and remove the glass stopper

F. Heat the flask over a Bunsen

burner, thus melting the butter and boiling the water. Watch the con-

^{*} U. S. Dept. of Agric., Bur. of Animal Ind., Circ. 100.

densation of the steam in the graduated part of the tube C, and do not allow the steam to get higher than the 15% mark. In case of continued foaming, allow the mixture to cool, add 2 cc. of the amyl reagent, and continue heating. After the water in the sample has boiled out, the temperature rises and the amyl reagent boils, driving the last traces of water and water-vapor from the flask and bottom of the stopper. Some of the amyl reagent is carried into the tube C with the steam, and some is boiled over after the water has been driven off. This amyl reagent in the tube is no disadvantage. When the mixture in the flask becomes a brown color and all the crackling noises in boiling cease, which usually requires 5 to 8 minutes, it is safe to conclude that all water has been driven from the flask.

Disconnect the flask A from the stopper B, place the glass stopper F in the tube C, giving it a turn to insure its being held firmly; invert the tube C, first being sure that the mouth of the small tube inside the bulb is held upwards, pour the water from the condensing jacket E, and remove the jacket. When the tube C is inverted, the water and reagent flow into the graduated part of the tube. To separate these and to get the last traces of water down into the graduated part, the tube C is held with the bulb in the palm of the hand, and the stoppered end away from the body, raised to a horizontal position, and swung at arm's length sharply downward to the side. This is repeated a number of times until the dividing line between the water and reagent is very distinct, and no reagent can be seen with the water or vice versa. The tube should then be held a short time with the stoppered end downward, and the amyl reagent in the bulb agitated in order to rinse down any adhering water.

The reading should not be taken until the tube and contents have cooled so little warmth is felt. When 10 grams of butter are used, the percentage is read directly at the lower meniscus.

With butter very low in moisture it may be desirable to use 15 grams, and with butter very high, 5 grams.

Fat.—This may be determined either directly or indirectly. For the direct determination, a weighed amount of the sample, from 2 to 3 grams, is first dried at 100° in sand or asbestos, contained in a thin and fragile round-bottomed evaporating-shell (Hoffmeister's Schälchen). If desired, the moisture may be determined in this connection by loss in weight after drying. The shell is afterwards inclosed in a piece of fat-free filter-paper, and crushed in pieces between the fingers in such a manner as to avoid loss. The pieces are gathered in a mass and folded together

in the filter-paper to form a packet of a size readily transferable to a Soxhlet extractor, in which the fat is removed in the usual manner and weighed, after drying, in a tared flask.

Or, the fat may be indirectly determined by subtracting the sum of the water, casein, and ash from 100.

Casein.—The residue from the determination of water by the A.O. A. C. method is stirred with petroleum ether until the fat is dissolved, and transferred to a tared Gooch crucible. After thorough washing with petroleum ether, the crucible is dried at 100°, cooled, and weighed, thus obtaining the casein and ash. The loss on ignition at a dull red heat represents the casein.

If desired nitrogen may be determined in the residue after removal of the fat with petroleum ether, and casein calculated from the nitrogen, using the factor 6.37.

Ash.—The residue left on the Gooch crucible after ignition, obtained as described in the preceding section is the ash. It consists largely of salt, which may be calculated from the percentage of chlorine determined by titration.

Milk Sugar and Lactic Acid compose most of the undetermined matter remaining after deducting from the total solids the sum of the fat, casein, and ash. Determine milk sugar, if desired, in an aqueous extract of the butter by Fehling's solution.

Determination of Salt.—In a tared dish or beaker weigh out about 5 grams of butter, taking a gram or so at a time from different parts of the sample. Add hot water to the weighed part, and after it has melted, the contents of the dish are poured into a separatory funnel, shaken and allowed to stand till the fat collects at the top, after which the underlying aqueous solution is drawn off into an Erlenmeyer flask, leaving the fat in the funnel bulb. Hot water is again added, and from ten to fifteen extractions are made, using about 20 cc. of water each time, all the water being collected in the Erlenmeyer flask.

A few drops of a solution of potassium chromate are then added for an indicator, and the sodium chloride volumetrically determined by a standard silver nitrate solution.

Salted butter contains from 0.5 to 6% of salt.

Examination of Butter Fat.—The butter fat is best obtained free from curd and salt by filtering when hot, the sample being best melted in a beaker on the water-bath. The water, with the curd and salt, will settle to the bottom. The clear fat is then filtered at a temperature not

exceeding 50° C., and subjected to such examination as may be desired to determine its purity.

U. S. Standard Butter Fat has a Reichert-Meissl number not less than 24 and a specific gravity not less than 0.905 at $\frac{40^{\circ}}{40^{\circ}}$ C.

ADULTERATION OF BUTTER.

The artificial coloring of butter is an art practiced for so many years, and is so far in accord with the popular demand, that it can hardly be considered as an adulteration. The most recent custom of adding preservatives other than salt to butter is, however, very properly considered in most localities as reprehensible, unless the character and amount of the preservative be made clear to the purchaser by a suitable label.

The most common and time-honored sophistication is the substitution in whole or in part of foreign fat, as in the case of oleomargarine, and, more recently, in the fraudulent sale of renovated or process butter for the freshly made article.

U. S. Standard Butter is butter containing not less than 82.5% of butter fat. By acts of Congress approved August 2, 1886, and May 9, 1902, butter may also contain added coloring matter.

ARTIFICIAL COLORING MATTER IN BUTTER.—Formerly carrot juice and annatto were used almost entirely as butter colors. The carrot furnished to the farmer a ready means of coloring his dairy butter, and its use was long in vogue for this purpose, before the commercial butter colors were available. Other vegetable colors, such as turmeric, marigold, saffron, and safflower, are said to have been used for this purpose, but, with the possible exception of turmeric, the writer is not aware of authentic cases in which they have been found in recent years. While annatto as a butter color is still in use, it is rapidly giving place to various oil-soluble, azo coaltar colors, which are admirably adapted to the purpose. All butter colors are now put on the market in solution in oil, usually cottonseed in this country and sesame in Europe.

Detection.—Martin* devised a general scheme, applicable for the detection of various colors in butter. His reagent consists of a mixture of 2 parts of carbon bisulphide with 15 parts of ethyl or methyl alcohol. 25 cc. of this solution are shaken with about 5 grams of the butter to be tested, and, after standing for some minutes, the mixture separates into two layers, of

^{*} Analyst, 12, p. 70.

which the lower consists of the fat in solution in the carbon bisulphide, while the upper is the alcohol, which dissolves out and is colored by the artificial dye employed. If saffron is present, the alcoholic extract will be colored green by nitric acid and red by hydrochloric acid and sugar.

Coal-tar dyes, if present, may be fixed on silk or wool by boiling bits of the fiber in the alcoholic extract, diluted with water and acidulated with hydrochloric acid.

Turmeric is to be suspected, if ammonia turns the alcoholic extract brown; marigold, if silver nitrate turns it black, and annatto, if on evaporating the alcoholic solution to dryness and applying to the residue a drop of concentrated sulphuric acid, a greenish-blue coloration is produced.

Turmeric is further tested for in the residue from the alcoholic extract as above obtained, by boiling the residue in a few cubic centimeters of a dilute solution of boric acid (or a solution of borax acidulated with hydrochloric acid), and soaking a strip of filter-paper therein. On drying the paper, if it assumes a cherry-red color, turning dark olive by dilute alkali, the presence of turmeric is assured.

Carrotin (the coloring matter of the carrot root) does not impart its color to the alcohol layer in Martin's test. Moore * has pointed out this exception, and shown that while with carrotin present the alcohol layer in Martin's test remains colorless, as in the case of uncolored butter, that when, however, a drop of very dilute ferric chloride is added, and the test-tube shaken, if carrotin be present, the alcohol will gradually absorb the yellow color from the butter. Care must be taken to avoid an excess of ferric chloride, as very little of this reagent will suffice.

Allen states that a butter color commercially known as "carrotin" consists in reality of 1 part of annatto in 4 parts of oil.

Detection of Annatto in Butter.—Treat 2 or 3 grams of the melted and filtered fat (freed from salt and water) with warm, dilute sodium hydroxide. After stirring, pour the mixture while warm upon a wet filter, using to advantage a hot funnel. If annatto is present, the filter will absorb the color, so that, when the fat is washed off by a gentle stream of water, the paper will be dyed straw color. It is well to pass the warm alkaline filtrate two or three times through the fat on the filter to insure removal of the color.

If, after drying the filter, the color turns pink on application of a drop of stannous chloride solution, annatto is assured.

^{*} Analyst, 11, p 163.

Detection of Coal-tar Colors in Butter.—Geisler's Method.*—A few drops of the clarified fat are spread out on a porcelain surface and a pinch of fullers' earth added. In the presence of various azo-colors, a pink to violet-red coloration will be produced in a few minutes. Some varieties of fullers' earth react much more readily with the azo-dyes than do others. In fact some do not respond at all. When once a satisfactory sample of this reagent is obtained, a large stock should be secured of the same variety.

Low's Method.†—A small amount of material to be tested is melted in a test-tube, an equal volume of a mixture of 1 part of concentrated sulphuric acid and 4 parts of glacial acetic acid are added, and the tube is heated nearly to the boiling-point, the contents being thoroughly mixed by shaking; the tubes are set aside, and after the acid solution has settled out it will have been colored wine-red in the presence of azo-color, while with pure butter fat, comparatively no color will be produced.

Doolittle's Method for Azo-colors and Annatto.‡—The melted sample is first filtered. Two test-tubes are taken and into each are poured about 2 grams of the filtered fat, which is dissolved in ether. Into one test-tube are poured 1 or 2 cc. of dilute hydrochloric acid, and into the other about the same volume of dilute potassium hydroxide solution. Both tubes are well shaken and allowed to stand. In the presence of azo-dye, the test-tube to which the acid has been added will show a pink to wine-red coloration, while the potash solution in the other tube will show no color. If annatto has been used, on the other hand, the potash solution will be colored yellow, while no color will be apparent in the acid solution.

Cornelison's Test for Artificial Colors.§—Melt 10 grams of the clear, dry fat, and shake well in a separatory funnel with 10 to 20 grams of 99.5% acetic acid. If the materials are too hot, the fat will dissolve, but at about 35° it separates quickly and almost completely. Draw off the clear acid, and after noting its color, test by adding to one portion of 5 cc. a few drops of concentrated nitric acid, and to another portion a few drops of concentrated sulphuric acid.

Natural yellow butter gives by this test a colorless extract, which remains colorless on adding nitric or sulphuric acid. The acid extracts of annatto, curcumin, and carrot are various shades of yellow, both before

^{*} Jour. Am. Chem. Soc., 20, 1898, p. 110.

[†] Ibid., 20, p. 889.

[‡] U. S. Dept. of Agric., Bur. of Chem., Bul. 65, p. 152.

[§] Jour. Am. Chem. Soc., 30, 1908, p. 1478.

and after addition of nitric acid, while with sulphuric acid they take on a pink coloration on standing, which in the case of curcumin is very decided. Soudan I and butter yellow give pink extracts, which remain pink on adding the stronger acids, while cerasine orange G, yellow O.B., yellow A.B. and certain other coal-tar dyes give extracts of various shades of yellow, which on treatment with the heavy acids in some cases remain colorless, but in others become pink, while the oil globule which separates remains colorless or takes on a pinkish color according to the dye.

PRESERVATIVES AND THEIR DETECTION.—Fresh or unsalted butter and renovated butter are often found with an added preservative, the one most commonly used for this purpose being the so-called "boric mixture" (borax and boric acid) already discussed under milk adulteration. Salted butter is occasionally, though not so often, found preserved. Other preservatives used in butter are formaldehyde, and salicylic and sulphurous acids. These latter are, however, rarely found.

Boric Acid.—This, if present, is best detected in the aqueous solution that settles to the bottom when butter is melted at the temperature of the boiling water bath, the supernatant fat being decanted off. Richmond* claims to be able to distinguish free boric acid from borax as follows: If on applying turmeric-paper directly to the aqueous liquid the paper turns red, the color being especially evident on drying, free boric acid is indicated. As a confirmatory test the reddened turmeric-paper is treated with dilute caustic alkali, whereupon it turns a dark olive-green if boric acid is present.

In the absence of a red color by the above test, or when this color is faint, the aqueous solution is acidified slightly with hydrochloric acid and the turmeric-paper applied as before. If borax be present to an appreciable extent, the red color will now be quite marked, even though not appearing before. In other words, testing with turmeric-paper without acidifying with hydrochloric acid shows, according to Richmond, a slight coloration due to the free acid alone, while the more intense color formed by first acidifying is due to the combined acid or borax.

Determination of Boric Acid.—Ten grams of the butter fat are weighed in a beaker and transferred with hot water to a separatory funnel in which the fat is extracted with 10 to 15 portions of hot water as described on page 534. The combined aqueous extract is evaporated to dryness in a platinum dish, the residue made alkaline, and ignited at

^{*} Dairy Chemistry, p. 254.

a dull red heat. Boil the ash with water, filter, and wash with hot water, keeping the volume of the filtrate under 60 cc. Make sure that the solution is perfectly neutral to methyl orange by treatment, if necessary, with sulphuric acid and tenth-normal alkali, add 30 cc. of glycerin, a few drops of the phenolphthalein indicator, make up to 100 cc., and titrate with tenth-normal sodium hydroxide according to Thompson's method (p. 823).

Butter being practically free from phosphates, the preliminary treatment for removing phosphoric acid in Thompson's method may be omitted.

Formaldehyde.—The aqueous solution from which the fat of the butter melted at low temperature has been poured off, is added to some milk previously found free from formaldehyde, and the test for the latter with hydrochloric acid and ferric chloride is tried directly in the milk.

Salicylic Acid.—Detection.—See method No. 2 for detection in milk, page 183.

Determination of Salicylic Acid.—Method of the Paris Municipal Laboratory.—Repeatedly exhaust 20 grams of butter in a separatory funnel with a solution of sodium bicarbonate, thus obtaining soluble sodium salicylate, if salicylic acid be present. Acidulate the aqueous extract with dilute sulphuric acid, and extract with ether. Evaporate the ether, and to the residue add a little mercuric nitrate, forming a precipitate nearly insoluble in water. Filter this off, wash the precipitate with water, and decompose into free salicylic acid with dilute sulphuric acid. Redissolve in ether, evaporate the solvent as before, and dry the residue at a temperature of 80° to 100°. Extract the residue with petroleum ether, dilute the ethereal liquid with an equal volume of 95% alcohol, and titrate with tenth-normal alkali, using phenolphthalein as an indicator.

1 cc. of tenth-normal alkali=0.0138 gram salicylic acid.

Sulphurous Acid.—The aqueous liquid, separated from the butter fat, is distilled, and the distillate treated with bromine water and barium chloride. A precipitate on the addition of the latter reagent indicates the presence of sulphurous acid or a sulphite in the butter.

Glucose in Butter.*—Crampton states that glucose has been found by him in butter intended for export to tropical countries, added to pre-

^{*} Jour. Am. Chem. Soc., 20, 1898, p. 201.

vent decomposition. In one sample made for export to Guadeloupe he found over 10% of glucose.

For its detection or estimation 10 grams of the sample are weighed out and transferred to a separatory funnel with hot water, and shaken out with successive portions of hot water. These are combined, and the aqueous extract made up to 250 cc. The reducing sugar may be determined by Fehling's solution or by polarization, using in the latter case alumina cream as a clarifier. While a slight reduction should be disregarded, any considerable reduction may be undoubtedly ascribed to glucose.

BUTTER "FILLED" WITH WATER.—Various preparations have been placed on the market to aid in incorporating water with butter. So called "black pepsin" has been used for this purpose. By churning the butter with water and a certain amount of the preparation in such a manner as to destroy the grain, it is possible to introduce two or three times the normal amount of water.

RENOVATED OR PROCESS BUTIER.

This product is also variously termed "boiled," "aerated," and "sterilized" butter. There are various modifications of the process of manufacture, but the object is to melt up and treat rancid butter in such a manner that for a time at least it is sweet. The following manner of treatment is typical, and shows in the main the necessary steps in carrying out the process, though details of manipulation vary in different localities.

The butter is melted in large tanks surrounded with hot water jackets at a temperature varying from 40° to 45° C. By this means the curd and brine settle to the bottom, whence they are drawn off, while the lighter particles rise to the top in the form of a froth or scum and are removed by skimming.

The clear butter fat is then, as a rule, removed to other jacketed tanks, and, while still in a molten condition, air is blown through it, which removes the disagreeable odors. The melted fat is then churned with an admixture of milk (more often skimmed) till a perfect emulsion is formed, after which it is rapidly chilled by running into ice cold water, with the result that it becomes granular in form. It is then drained and "ripened" for some hours, after which it is worked free from excess of milk and water, salted, and packed.

Under some state laws this product, to be legally sold, must conform to rules of labeling as strict as those prescribed for oleomargarine. In other localities it may be sold with impunity. Not infrequently it is sold as choice creamery butter, and sometimes at the same price.

U. S. Standard Renovated or Process Butter should contain not more than 16% of water, and at least 82.5% of butter fat.

OLEOMARGARINE.

According to the U. S. revenue laws, artificial butter composed wholly or in part of fat other than butter fat must be branded oleomargarine. The name butterine, although used in advertising matter, does not have the sanction of the government. The product is commonly known in England as margarine. As a rule the oleomargarine of commerce is composed of refined oleo oil, usually churned up with neutral lard, milk, and a small amount of pure butter, the whole being salted and sometimes colored to resemble butter. Cottonseed oil and other vegetable oils are also used to some extent.

Oleo oil is prepared from the fat of beef cattle somewhat as follows:* Immediately after the animals are killed the fresh intestinal and caul fat are removed and placed in tanks of water at a temperature of about 80° F. From this water they are transferred to other tanks of cold water and chilled until all animal heat is removed. The fat is then cut or hashed into small pieces and melted at about 150° F. in jacketed steam kettles, until the clear oil is separated from the connective tissue.

This oil is then drawn off into vats, which, on account of the appearance of the oil on cooling, are called graining or seeding vats, where it is allowed to stand for twenty-four hours or more at a temperature of about 85° F. From these vats the semi-solid emulsion of oil and stearin is dipped into cloths, which are folded and placed in a press between sheets of metal and subjected to powerful pressure. By this means the oil is separated from the stearin, and is drawn into casks for export or for manufacture into oleomargarine. Large quantities are annually exported to Holland, where oleomargarine is manufactured, and either sold for consumption in that country, or re-exported to other countries in Europe.

The oleo oil thus expressed is a mixture of olein and palmitin. When first prepared, it is a clear amber-colored fluid, free from odor or fatty taste. It is packed in tierces, and, when opened at ordinary temperature, is a light-yellow solid.

The further process of manufacture of oleomargarine consists in

^{*} Report on Oleomargarine, Its Manufacture and Sale, 19th An. Report, Mass. St. Bd. of Health, 1887.

the main of mixing the oleo oil as above obtained with varying proportions of neutral lard, milk, and genuine butter, with or without added coloring matter, and churning the mixture at a temperature above the melting-point of the fats, the neutral lard having previously been cured for at least forty-eight hours in salt brine. Occasionally small quantities of other vegetable oils, as cottonseed, peanut, or sesame, are included in the above mixture. After the churning, the whole mass is cooled by contact with ice water. The chilled mass is drained, and afterwards salted, worked, and given much the same treatment as butter.

The composition of commercial oleomargarine varies between the following limits:

Oleo oil	20	to	25%
Neutral lard	40	"	45%
Butter	10	"	25%
Milk, cream, salt, etc.	5	"	30%

Coloring of Oleomargarine.—The artificial coloring matters employed are the same as in the case of butter, and are similarly tested for.

In many states oleomargarine cannot be legally sold when colored to resemble butter. Under other state laws coloring matter is allowable. The federal law and most state laws prescribe the most rigid rules for marking packages containing oleomargarine, with a view to affording the utmost protection to the producer of butter against the fraudulent substitution therefor.

Crampton and Simon's Tests for Palm Oil.*—So called "butter oils," consisting of cottonseed oil to which has been added 2 to 5 per cent of palm oil are used to color oleomargarine. The following tests serve for the detection of palm oil.

Preparation of Sample.—The sample should be kept in a cool, dark place until tested, as exposure to air and light, or the presence of water, alcohol, ether or similar reagents interfere with the tests. Immediately before testing, the sample is filtered as quickly as possible at a temperature not exceeding 70° C.

First Method.—Dissolve 100 cc. of the fat in 300 cc. of petroleum ether, and shake out with 50 cc. of 0.5% potassium hydroxide. Draw off the watery layer, make distinctly acid with hydrochloric acid, and shake out

^{*} Jour. Am. Chem. Soc., 27, 1905, p. 270.

with 10 cc. of colorless C. P. carbon tetrachloride. Separate the carbon tetrachloride solution, transfer a portion to a porcelain crucible, add 2 cc. of a mixture of one part of colorless, crystallized C. P. phenot and 2 parts of carbon tetrachloride, then 5 drops of hydrobromic acid (sp. gr. 1.19), and mix by gentle agitation.*

The almost immediate development of a bluish-green color is indicative of palm oil.

Second Method.—Shake 10 cc. of the melted and filtered fat with an equal volume of colorless C. P. acetic anhydride, add one drop of sulphuric acid (sp. gr. 1.53), and shake a few seconds longer.†

If palm oil be present, the lower layers on settling out will be found to be colored blue with a tint of green. The color in this as in the preceeding test is transient.

Of the edible oils only sesame and mustard oils give a similar color reaction. Sesame oil, after repeated extractions with alcohol, will not give the blue color, but cottonseed oil containing as little as 1% of palm oil still responds to the test.

Adulterants of Oleomargarine.—This product is liable to adulteration not only by the use of inferior and unwholesome fat, but by the admixture in some cases of paraffin.‡ This sophistication is made manifest, if an appreciable amount of the adulterant has been used, by the high melting-point and the low saponification number, as well as by the low specific gravity. If a clear saponification is impossible under ordinary conditions, paraffin is to be suspected. It may be separated and quantitatively determined as described on p. 510.

Healthfulness of Oleomargarine.—Under the directions of the Massachusetts Board of Health, a large number of artificial digestion experiments were made to show the relative nutritive value of butter and oleomargarine, and at the same time the wholesomeness of oleomargarine as a food was carefully investigated. The general conclusions reached were that, when comparing the best grades of both products, there is little if any difference between butter and oleomargarine on grounds of digestibility, while a good oleomargarine is much to be preferred to a

^{*} Halphen uses a similar reagent to detect rosin oil in mineral oil. Jour. Soc. Chem. Ind., 21, 1902, p. 1474.

[†] The reagents are the same as used in the Liebermann-Storch test for rosin oil.

[‡] Geissier, Jour. Am. Chem. Soc., 21, 1899, p. 605.

¹⁹th An. Report, Mass. State Board of Health, 1887, p. 248.

poor butter from a nutritive standpoint. As to its wholesomeness, a large number of experts consulted were unanimous in expressing their favorable opinions of oleomargarine as a healthful article of food.

When sold on its own basis in accordance with the law, it forms an excellent cheap substitute for butter. It is only when fraudulently sold as butter or in violation of the various state and federal laws, that it comes within the province of the health authorities to condemn it, and, unfortunately, by reason of its close resemblance to the dairy product the temptation to sell it for what it is not is always great.

Distinguishing Oleomargarine from Butter.—The two products, made up as they are of mixtures of the same fats, and differing for the most part only in the percentage composition of these fats, show many properties in common. For instance, the melting-point is so nearly the same for both products as to be of no use as a distinguishing indication. Other physical characteristics, as of taste and smell, are very similar in both products, except in the hands of the expert. The microscope is of limited value, except in so far as it indicates that the fat has first been melted and afterwards solidified.

From the fact that oleo oil and neutral lard form by far the larger portion of the mixture known as oleomargarine, the glycerides that make up the fat of the latter are chiefly those of the insoluble fatty acids, stearic, oleic, and palmitic. The percentage of volatile fatty acids present in oleomargarine is very small, and the presence of these volatile acids is due entirely to the admixture of butter which it contains. This furnishes the most ready means of distinguishing chemically between the two products, and, as indicated by the Reichert number, is the chief reliance of the analyst for court evidence.

Incidentally, as will be seen by the accompanying table, the refrac-

CONSTANTS OF BUTTER FAT AND OLEOMARGARINE.

	Specific Grav- ity at 100°.	Water.	Insoluble Acids.	Soluble Acids.	NaCi.	Albuminoids.	Curd.	Koettstorfer's Equivalent.	Reichert Number.*	Butyro-re- fractometer Reading.	Temperature.
Minimum Oleomargarine	.867	31.55 4.44 11.69 9.34	85.63†	3.00\$	0.00‡	.175‡	3.10 0.49 0.74 0.63	233 2225 203 192	15.8† 12.4† 5.5† 0.5†	44.81	35° 35° 35°

Number of cubic centimeters N/10 alkali neutralizing volatile acids in 2.5 grams fat. From analyses made in Mass. State Board of Health laboratory. From analyses made in laboratory of U. S. Dept. of Agric., Bur. of Chem. From analyses by A. H. Allen.

tometer reading, the iodine number, the saponification equivalent, and the specific gravity are all useful constants in indicating points of difference between the two fats, it being understood that in oleomargarine, as in butter, the fat for examination is melted and separated by filtration or otherwise from the curd, salt, and other constituents.

The constants for varying mixtures of butter with foreign fat as found by Villiers and Collin* are tabulated below.

Odor and Taste.—It is easy with a little practice to become so accustomed to the odor and taste of oleomargarine, as to be able to pass judgment with considerable confidence by these senses alone, whether a sample in question is oleomargarine or butter. The distinction is rendered more apparent by melting a portion of the sample on the waterbath. If the product is butter, either fresh or renovated, the butyric odor of the melted fat is very characteristic, while the melted oleomargarine not only is lacking in the butyric odor (a negative property), but possesses a distinctive "meaty" smell peculiar to itself, which, while not unpleasant, is unmistakable. The flavor of oleomargarine to one experienced in distinguishing between the two products is very apparent. This flavor, slight though it is, might be compared to that of cooked meat.

	Hehner's Number.	Soluble Acids.	Koettstorfer's Equivalent.	Volatile Acids.
Pure butter	88	5	224	26 .
Butter, 95%; foreign fat, 5%	88.35	4.8	222.6	24-7
" 90% " " 10%	88.70	4.5	221.2	23.4
" 85% " " 15%	80.05	4-3	219.8	22.2
" 80% " " 20%	89.40	4.1	218.4	20.9
" 75% " " 25%	89.75	3-9	217	19.6
" 70% " " 30%	90.10	3. 6	215.6	1 8. 3
" 65% " " 35% ····	90-45	3-4	214.8	17. Ĭ
" 60% " " 40%	90.80	3.2	212.8	15.8
55% 45% 45% 	91.15	3	211.4	14.5
" 50% " " 50%	91.50	2.7	210	13.2
45% " 55%	ór.85	2.5	208.6	12
" 40% " " 60%	92.20	2.3	207.2	10.7
35% " " 65%	92.55	2. I	205.8	9-4
" 30% " " 70%	92.90	1.8	204.4	8. i
" 25% " " 75%	93.25	1.6	203	6.9
" 20% " " 80%	93.60	1.4	201.6	5.6
" 15% " " 85%	93.95	1.2	200.2	4-3
" 10% " " 90%	94.30	0.9	198.8	3
" 15% " " 85% " 10% " " 90% " 5% " " 95%	94.65	0.7	197.4	ĭ.8
Foreign fat.	95	0.5	196	0.5

^{*} Les Substances Alimentaires, p. 731.

DISTINGUISHING BETWEEN BUTTER, PROCESS BUTTER, AND OLEOMARGARINE.

With the increased occurrence in the market of the commercial product known as "process" butter, especially in localities where its sale is restricted or regulated by law, it becomes incumbent on the analyst to distinguish it from the other products which it resembles.

As a rule, the tests, chiefly physical, that are applied on the edible product as a whole (i.e., without separation of the curd, salt, etc.), such as the foam test, the milk test, the microscopical examination, and the appearance of the melted sample, distinguish broadly between pure fresh butter on the one hand, and oleomargarine on the other. In other words, although there are those skilled in making the above tests who claim to be and doubtless are able to note distinguishing features between oleomargarine and process butter, yet these two products respond alike, though perhaps in varying degrees, to these tests, and are classed together as distinguished from pure butter.

On the other hand, such tests as depend upon the refractometer, the Reichert number, and, indeed, all the so-called chemical constants, which are applied to the separated fat, freed from other substances, will serve to distinguish between oleomargarine and butter, whether "process" butter or otherwise, since the "processing" or "renovating" of butter does not change the character of its fat sufficiently to materially alter these constants.

It is best, therefore, for purposes of routine preliminary separation to submit all samples to the "foam" test and to examine them by the butyro-refractometer.* These tests alone, which are very quickly and readily applied, will rarely fail to separate into the three classes, butter, process butter, and oleomargarine, the products under examination, after which such confirmatory tests as are desired are made on adulterated samples.

The Butyro-refractometer.—This instrument, as its name implies was primarily intended by Zeiss for the examination of butter, and, while its use has been extended for work with other fats and oils, its construction is such as to show particularly a distinction between butter and oleomargarine by the appearance of the critical line of the fat. This mode of differentiation is due to the peculiar construction of the double

^{*} Out of the large number of samples of butter and oleomargarine examined on the butyro-refractometer in the author's laboratory during eight years, he has never found a single instance where the instrument failed to show the difference between the two products.

prism, which shows differences of dispersive power by different appearances of the critical line. The prisms are so constructed that the critical line of pure butter is colorless, while margarine and artificial butter, which have greater dispersive powers than natural butter, show a blue-colored critical line. But anomalies in the color, both with pure butter and mixtures, are more or less observable, which render it impossible to draw a sharp line between adulterated and genuine butter. The appearance of a blue fringe may, however, be a useful factor in cases of suspected adulteration.

The following particulars respecting the application of the refractometer for analysis of butter are contained in a paper of Dr. R. Wollny of Kiel,* who assisted in the construction of the instrument. The readings of the refractive indices of a large number of butter samples taken at 25° C. by Dr. Wollny have been directly reduced to scale divisions and yield the following equivalents:

```
Natural butter...(1.4590-1.4620):49.5-54.0 scale divisions
Oleomargarine...(1.4650-1.4700):58.6-66.4 ""

Mixtures (artificial butter) ......(1.4620-1.4690):54.0-64.8 ""
```

Limit of Scale Reading for Pure Butter.—Whenever in the refractometric examination of butter at a temperature of 25° C. higher values than 54.0 are found for the critical line, these samples will, according to Wollny, by chemical analysis always be found to be adulterated; but with all samples in which the value for the position of the critical line does not reach 54.0 chemical analysis may be dispensed with, and the samples may be pronounced to be pure butter. Wollny suggests, as a means of removing all chances of adulterated butter escaping detection, that the above limit be placed still lower, and that all samples exhibiting values exceeding 52.5 (at a temperature of 25° C.) be set aside for chemical analysis.

In calculating the position of the critical line for other temperatures than 25° C. allow per 1° C. variation of temperature a mean value or

^{*} Dr. R. Wollny, Schlussbericht über die Butteruntersuchungsfrage, Milchwirthschaftlicher Verein, Korrespondenzblatt, No. 39, 1891, p. 15.

Older papers on butter tests by refraction of light will be found in: Mueller, Rep. d. anal. Chemie, 1886, pp. 346, 366. Skalweit, Milchzeitung, 1886, 15, p. 462. Wollny, Ueber die Kunstbutterfrage, Leipzig, 1887, p. 50.

o.55 scale division.* The following table, which has been compiled in this manner, shows the values corresponding to various temperatures, each value being the upper limit of scale divisions admissible in pure butter:

Temper-	Scale	Temper-	Scale	Temper-	Scale	Temper-	Scale
ature.	Division.	ature.	Division.	ature.	Division.	ature.	Division.
45°	41.5	40°	44.2	35°	47.0	30°	49.8
44°	42.0	39°	44.8	34°	47.5	29°	50.3
43°	42.6	38°	45.3	33°	48.1	28°	50.8
42°	43.1	37°	45.9	32°	48.6	27°	51.4
41°	43.7	36°	46.4	31°	49.2	26°	51.9
40°	44.2	35°	47.0	30°	49.8	25°	52.5

If, therefore, at any temperature between 45° and 25° values be found for the critical line which are less than the values corresponding to the same temperature according to the table, the sample of butter may safely be pronounced to be natural, i.e., unadulterated butter. If the reading shows higher numbers for the critical line, the sample should be reserved for chemical analysis.

Note.—Dr. Eichel of Metz has suggested that instead of comparing the scale divisions at the same temperature, the position of the critical line may be determined at the moment when the butter begins to set. In this case he gives fifty-four as the highest admissible number for the critical line of pure butter.

No sharp distinction is apparent between pure and renovated butter on the refractometer.

Special Thermometer for the Butyro-refractometer.—Instead of employing the ordinary thermometer, as shown in Fig. 36, a special thermometer (Fig. 101) has been devised for work both with butter and with lard. This instrument has two scales, arranged side by side, one for butter and one for lard, each of which indicates at once the highest allowable reading for the pure fat, corresponding to the temperature at which the observation is made, which, however, need not be noted.

If the scale reading of the instrument, as observed through the telescope, differs materially from the reading of the special thermometer, the fat under examination is undoubtedly adulterated, or, in the case of butter, a higher reading indicates oleomargarine. The special thermometer thus indicates the highest permissible number for pure butter.

^{*} With natural butter this number is, as a rule, somewhat less (0.53), with oleomargarine a little greater (0.56).

The Reichert or Reichert-Meiasl Number* is by far the most important single determination in establishing proof of the character of the sample, whether butter or oleomargarine, for evidence in court, and in such cases this determination is indispensable. The result is conclusive, excepting in those rare instances where the admixture of foreign fat is so small as to cause the Reichert number to approximate that of pure butter. In common instances of creamery butter and commercial oleomargarine the Reichert number shows a very marked distinction (see table, p. 544).

It is difficult to fix a minimum figure below which, in doubtful cases, a sample may be pronounced impure by reason of admixture with foreign fat. In general, however, a Reichert number under 10 would be almost sure to show adulteration, though instances are on record where butter of known purity has shown a Reichert number even lower than this. It is in fact rare that pure butter has a Reichert number under 12.

Stebbins † gives the maximum, minimum, and average of the Reichert number obtained by him on 317 samples of unadulterated butter, some of which were of low grade, Fig. 101. - Special as follows: Maximum, 18.2; Minimum, 11.2; Average, 14.7. Butyro-refrac-

As a rule little difference is apparent between pure and "renovated" samples as regards their Reichert number.

Butyro-refractometer Thermometer for Butter and Lard.

Vieth has shown that the Reichert number of butter is generally a trifle lower after it becomes rancid.

Specific Gravity.—Skalweit has shown that the specific gravity of butter and oleomargarine relative to each other varies with the temperature at which it is taken, the difference between the two growing less and less as the temperature increases above 35°. The greatest variation being at 35°, he recommends this temperature as the best at which to make the determination.

The Foam Test, also known as the "boiling" or "spoon" test.‡ This, though originally intended as a household test, is in reality one of

^{*} The writer prefers to carry out this process on 2.5 grams of the butter fat, expressing thus the Reichert number, this being practically half the Reichert-Meissl number, which is based on the use of 5 grams.

[†] Jour. Am Chem. Soc., 21, 1899, p. 939.

[‡] Farmer's Bulletin, No. 131.

the very best laboratory methods of separating pure butter samples from renovated butter and oleomargarine. A small lump of the sample (from 3 to 5 grams) is heated in a large spoon over a Bunsen flame, turned very low, stirring constantly during the heating. Genuine butter, under these conditions, will boil quietly, but with the production of considerable froth or foam, which will often swell up over the sides of the spoon, when, just after boiling, the latter is raised from the flame. Renovated butter or oleomargarine, under this treatment, will bump and sputter noisily like hot grease containing water, but will not foam.* Another point of difference is that on removing the spoon from the flame and observing the character of the curdy particles, in the case of genuine butter these particles of curd will be very small and finely divided in the melted fat, being indeed hardly perceptible, while with oleomargarine and renovated butter, the curd will gather in somewhat large masses or lumps.

The test may be carried out in a test-tube if desired.

The Waterhouse or Milk Test.†—This test is based on the assumption that butter fat, which is in itself exclusively the product of milk, will mingle intimately with the milk when added thereto in a melted condition and cooled therein, whereas oleomargarine, being foreign to milk fat, will, under like conditions, refuse to diffuse itself naturally in milk as a medium.

About 50 cc. of well-mixed sweet milk are heated nearly to boiling in a beaker, and from 5 to 10 grams of the fat sample are added. The mixture is then stirred, preferably with a small wooden stick, until the fat is melted. The beaker is then placed in a dish of ice cold water, and the stirring continued till the fat reaches the solidifying-point, at which period, if the sample is oleomargarine, the fat can readily be collected by the stirrer into one lump or clot, but, if butter, it cannot be so collected, but remains in a granulated condition, distributed through the milk in small particles. It is not necessary to keep up the stirring through the entire term of cooling, but to begin stirring before the fat starts to solidify, which should require from ten to fifteen minutes after the mixture is placed in cold water.

This test, if carefully carried out, shows a marked distinction between butter, whether pure or renovated, and oleomargarine. Under certain conditions, as when the cooling is too rapid, samples of renovated butter

^{*} A very slight foam is sometimes observable with occasional renovated samples, but nothing like the abundant amount produced by the genuine product.

[†] Parsons, Jour. Am. Chem. Soc., 23, 1901, p. 200.

fat will sometimes show a slight tendency to clot together as in the case of oleomargarine, but to no such extent as the latter.

The author's experience with this test has shown it to be very reliable not only in identifying oleomargarine from butter, but in nearly every case renovated butter can be distinguished from genuine. As a rule, genuine butter fat, even after cooling to the solidifying-point, shows the greatest tendency to emulsionize with the milk when stirred, without adhering to the wooden rod, and is slow to come to the surface when the stirring is stopped. Renovated butter fat, when stirred in the cold milk, almost instantly gathers in a film on the surface of the milk when the stirring is stopped, without emulsionizing. It does not clot together like oleomargarine, but it tends to adhere to the wooden rod.

Patrick* recommends the use of skimmed or partially skimmed milk, and heats to the boiling-point after the fat has been introduced into the hot milk.

Examination of the Curd.—The curd of genuine butter is made up largely of such of the milk proteins as are insoluble in water and hence pass into the cream when separated. These proteins form a gelatinous mass in the butter, readily clotting together when the fat is melted. On the other hand, the curd of process butter, which is, as it were, artificially derived from the entire or skim milk used in its manufacture (in order to replace the natural curd which has been removed in the "purifying" process), differs from the proteins of cream in that it is granular and flaky, consisting chiefly of coagulated casein. Hence the distinction noted as to the appearance of the curd in the foam test.

For the same reason, if beakers containing pure and renovated butter are melted on the water-bath, the curd of the pure sample will settle at once, or in a very few minutes, to the bottom after melting, leaving a comparatively clear supernatant fat. The renovated sample will nearly always fail to settle out clear, even after standing on the water-bath for half an hour or more, but will still be cloudy throughout the mass, due to particles of non-cohesive, floating curd.

In the case of oleomargarine, the curd of which is composed partly of pure butter curd (from cream proteins) and partly of the proteins of the milk with which it is churned, the cloudiness of the fat on melting depends on the relative proportion of milk proteins, and in general is not especially characteristic.

^{*} Farmer's Bulletin, No. 131.

Identification of the Source of the Curd.*—Half fill a small beaker with the sample and melt on the water-bath. Decant as much as possible of the fat and pour the rest, consisting largely of the water, salt, and curd, upon a wet filter. Acidify the filtrate, which contains the salt and soluble proteins, with acetic acid and boil. If the sample is pure butter, only a slight milkiness is found, indicating absence of albumins, whereas, in the case of process butter, a white, flocculent albuminous precipitate is produced.

Apply to the filtrate also Liebermann's test for albumin; i.e., add strong hydrochloric acid. If a violet coloration is produced, the sample is presumably "process" butter.

Microscopical Examination of Butter.—Considerable information may in general be gained by an examination of the sample under ordinary light and with a rather low power, say from 120 to 150 diameters. For examination in this way a bit of the sample on the edge of a knife blade is placed on the glass slide, and simply pressed lightly into a thin film by the cover-glass. A very characteristic difference between genuine and renovated butter is at once seen in the relative opacity of the fields. The fat film, in the case of fresh, pure butter, is much more transparent than that of the renovated. Again, the curd is so finely divided throughout the mass of genuine butter fat that the field is much more even than that of the renovated, wherein often large and opaque patches of curd are frequently distributed throughout the field.

When a renovated butter sample, mounted as above, is viewed by reflected light, for which purpose the microscope mirror is turned so as not to transmit light through the instrument, one sees a very dark and scarcely perceptible field; but the opaque patches of curd above referred to are strikingly apparent as white masses against a dark background.

With Polarized Light.—It has already been stated that the microscope is useful in showing whether or not a fat has been melted, the crystalline structure of the fat once melted and afterward cooled being rendered apparent, especially when viewed by polarized light. This fact has long been known and put to practical use in the identification microscopically of butter and oleomargarine.†

When viewed by polarized light between crossed Nicols under a low magnification, pure butter not previously melted should show no

^{*} Hess and Doolittle, Jour. Am. Chem. Soc., 22, 1900, p. 151.

[†] Hummel, ibid., 22, p. 327; Crampton, loc. cit., supra, p. 703.

crystalline structure, being uniformly bright throughout, and, if the selenite plate be used, should present an evenly colored field, entirely devoid of fat crystals. On the other hand, with process butter or oleomargarine, both of which have been melted and subsequently cooled, the crystalline structure should be marked, showing with polarized light a more or less mottled appearance, and a play of colors with the selenice.

Various conditions enter in to affect the reliability of the polarized light test. It is nearly always possible in cold weather to observe these distinctions in practice, as above described, in a sharp and striking manner. Figs. 269, 270, and 271, Pl. XXXVIII, show typical fields of the three products with crossed Nicols and selenite plate. The appearance of pure butter is perfectly blank, while oleomargarine presents a much more mottled appearance than renovated butter. Such well-defined points of variation as are shown in Plate XXXVIII are not always to be seen in practice, even in the hands of an expert. Pure butter sometimes exhibits a somewhat mottled field, due to a slight crystallization at some period in its history. In the summer-time, for instance, when butter melts so easily at ordinary temperature, these distinctions between pure and adulterated samples as shown by polarized light are by no means as satisfactory as in the winter.

Great care should be taken on this account, on the part of the collector of samples as well as the analyst, to keep the sample from melting under ordinary conditions before it is examined.

Hess and Doolittle's Method of Examining the Curd.*—A convenient portion of the sample of suspected butter is melted in a beaker, as much of the fat as possible is decanted off, and the remaining curd, washed free from fat with ether, is poured out on a glass plate and dried. A sample of pure butter is treated in like manner by way of comparison. When examined under a very low magnification of from 3 to 6 diameters, the curd from the pure sample will be seen to be non-granular and amorphous in appearance, while, in the case of renovated butter, the curd will appear very coarse grained and mottled.

Zega's Test for Oleomargarine.†—A portion of the filtered fat is poured into a test-tube and kept for two minutes in a boiling water-bath. I cc. of this fat is then measured with a hot pipette into a 50-cc. tube containing 20 cc. of a mixture of 6 parts ether, 4 parts alcohol, and I part glacial acetic acid. The tube is stoppered, shaken well, and cooled in

^{*} Jour. Am. Chem. Soc., 22, 1900, p. 151.

[†] Chem. Zeit., 1899, 23, 312; Abs. Analyst, 24, p. 206.

water at 15° to 18° C. In the case of pure butter fat, the solution remains clear for some time, a slight deposit being apparent only after standing an hour or more. With oleomargarine, a deposit is evident in a very short time, and in ten minutes a heavy precipitate comes down. With 10% of oleomargarine in butter, a separation occurs in about fifteen minutes. When a few solid particles have separated out, they are withdrawn and examined under the microscope. With genuine butter, long narrow rods appear, sometimes pointed at the ends, often bent, and grouped as a rule centrally in star-shaped bundles. Oleomargarine presents an appearance of bundles of fine needles, closely packed to form masses frequently resembling sheaves and dumb bells in shape.

Identification of Various Oils and Fats. — Cottonseed oil may be recognized, if present in butter or its substitutes, by the Halphen test, and sesame oil by the Baudouin test. Peanut oil is tested for by the Bellier or Renard test.

Cocoanut oil is sometimes said to be present in butter substitutes. It has a higher Reichert number than most adulterants, and hence a larger admixture of this than of other foreign fats could be used, without lowering the Reichert number of the whole below the allowable limits of pure butter. Its presence would, however, be rendered apparent by the low iodine and refractometer numbers and the high Polenske number.

LARD.

Nature and Composition.—Lard is the fat of hogs, separated by heat from the scraps or containing tissues. The choicest or highest grade of lard is known as leaf lard, and is derived from the fat which surrounds the kidneys. A comparatively small part of the lard of commerce is, however, strictly speaking, pure leaf lard. Most of it is derived from the whole fat of the animal by rendering, by the aid of steam under pressure, either in open kettle or in closed tanks, the former being used more often for rendering lard on a small scale, and the latter being the most common commercial method.

Next to the leaf, the fat from the hog's back is considered the bos in quality, after which is graded, in the order named, the fat from the head, the region of the heart, and the small intestines, the last two grades constituting what is commonly known as "trimmings."

Good lard is white and granular, having the consistency of salve. It has an agreeable, characteristic odor and taste.

The leaf or kidney fat furnishes also the source of the so-called neutral lard, already mentioned as an ingredient of oleomargarine. The leaf:

being first chilled and finely ground, is placed in the kettle and rendered at a temperature of from 40° to 50° C., at which heat only a portion of the lard separates. This portion is, while melted, washed with water containing salt or dilute acid, and forms the neutral lard, a product almost entirely free from odor. The remainder of the leaf is then transferred to the closed tank and subjected for some hours to steam under pressure at a temperature of 230° to 290° F., the resulting lard being graded as pure leaf lard.

The composition of the mixed fatty acids of lard is thus calculated by Twitchell:

Linoleic acid	10.06%
Oleic acid	49-39%
Solid acids (by difference), stearic and palmitic	48.55%

Lewkowitsch* gives the following constants for American lards made from fat from different parts of the animal:

	Specific Gravity		Maumené	Melting-po mann's †	Refractive Index.	
Fat from	at 100° C. (Water at 15° C. = 1.)	Iodine Value.	Number at	Temp. C. of Incipient Fusion.	Melted to a Clear Drop.	Butyro- refractom- eter at 40° C.
Head	o.8637 o.8629	66.2 66.6	33	24 24	44.8 44.8	52.6 52.5
Back	0.8631 0.8611 0.8621 0.8616	65.0 61.5 65.0	34 37 35	24 28.5 28.5	45.0 48.5 48.5	52.0 52.4 51.8
Leaf	o.8637 o.8615	65.1 62.2 59.0	38	31.5 -26 29	46 45 44	51.9 51.4 50.2
FootHam	0.8700 0.8589 0.8641 0.8615 0.8628	63.0 68.8 68.4 66.6 68.3	30 38 	28.5 24 26 26 26 26	44-5 40 45 44 44-5	52.0 44.8 51.9 51.9 53.0
Ham (German)	0.8597	55.0	30	32	46	49.2

Lard Oil.—This oil is obtained by subjecting lard contained in woolen bags to hydraulic pressure in the cold. The lard oil (chiefly olein) thus expressed constitutes nearly 60% of the whole, and the residue is known as lard stearin.

Lard oil is a thin fluid, pale yellow in color, and with varying specific

^{*}Oils, Fats, and Waxes, 1904, p. 781.

[†] Bensemann distinguishes between the temperature at which the fat begins to liquefy and that at which it becomes completely transparent.

gravity, due to varying conditions of pressure and temperature. It has a pleasant, though somewhat bland taste, and is used to some extent as an edible oil. It is used in France as an adulterant of olive oil, and with the Maumené, elaïdin, and nitric acid tests, it behaves much like olive oil.

According to the U. S. Pharmacopæia, the specific gravity of lard oil should be from 0.910 to 0.920 at 15° C.

At a temperature a little below 10° C. it should form a semi-solid white mass.

When it is brought in contact with concentrated sulphuric acid, a dark reddish-brown color should instantly be produced.

Lard oil should not respond to the Bechi test for cottonseed oil.

If 5 cc. of the oil, contained in a small flask, be mixed with a solution of 2 grams of potassium hydroxide in 2 cc. of water, then 5 cc. of alcohol added, and the mixture heated for about five minutes on a water-bath with occasional agitation, a perfectly clear and complete solution should be formed, which, on dilution with water to the volume of 50 cc., should form a transparent, light-yellow liquid, without the separation of an oily layer (absence of appreciable quantities of paraffin oils).

Adulterants of lard oil are cottonseed and corn oils.

Compound Lard.—The article so extensively made and sold under this name is a mixture consisting usually of lard stearin, beef stearin, and cottonseed oil. Sometimes no lard whatever is present, but only a mixture of beef and cottonseed stearins.

Lard stearin is the residue left in the cloths after the lard oil has been removed by pressure (p. 555).

Beef stearin is, similarly, the residue from which oleo oil has been expressed (p. 541). The cottonseed oil used is highly refined, and finally decolorized by mixing with fullers' earth and filtering.

U. S. Standards.—Standard Lard and Standard Leaf Lard are lard and leaf lard respectively, free from rancidity, containing not more than 1% of substances other than fatty acids, not fat, necessarily incorporated therewith in the process of rendering, and standard leaf lard has an iodine number not greater than 60.

Adulteration of Lard.—The mixture known as "compound lard" is quite commonly fraudulently sold for pure lard. Indeed, the adulterants of lard usually met with are cottonseed oil or stearin and beef stearin. Other oils said to have been used as adulterants are peanut, sesame, corn, and cocoanut. Formerly water was incorporated into

the fat to such an extent as to materially cheapen it, but this sophistication is now rare. Moisture is determined as in the case of butter.

The Butyro-refractometer Reading.—The refracting degree of cotton-seed oil on the butyro-refractometer is about 8.9 in excess of the standard refraction of lard, while that of beef tallow is about 3.8 less than the standard. If, therefore, the refractometer reading is unusually low, the presence of beef stearin is to be suspected; if unusually high, cottonseed oil should be looked for. A mixture of the two adulterants with pure lard such as is found in "compound lard," may sometimes, though not often, be found to give refractometric readings within the limits of pure lard.

Detection of Foreign Oils.—Cottonseed oil is best detected by the Halphen test. A very slight color reaction should not be taken as proof positive of the admixture of cottonseed oil, since it has been found that the fat of hogs fed on cottonseed meal gives a slight reaction with both the Bechi and the Halphen tests. Sesame and peanut oils are detected by their special tests. Corn oil is indicated by the abnormally high refractometric reading and iodine number, cocoanut oil by the high Reichert number, the high saponification equivalent, and especially the high Polenske number.

Beef stearin is difficult to identify chemically, but is usually distinguished by a microscopical examination of the fat after crystallization as follows:

The Microscopical Examination of Lard.—From 2 to 5 grams of the fat are dissolved in 10 to 20 cc. of ether * in a test-tube, and the solution allowed to stand 12 hours or over night at about 20° C., the testtube being loosely stoppered with cotton. The crystals obtained vary considerably with the condition of heat, amount of solvent, rate of crystallization, etc., so that the operator had best vary these conditions till he is satisfied that the best possible results have been obtained. It is often advantageous to separate the crystals first obtained by filtration from the mother liquor, and to redissolve in ether and recrystallize in a second test-tube. The crystals formed at the bottom of the test-tube are, for the purpose of thus purifying, separated from the mother liquor by filtration through a small filter, and the precipitate washed several The washing with ether should not be continued times with ether. so long that the crystals are perfectly freed from mother liquor and olein, for in this case they are so dry and pulverulent as to require a mountant when on the slide for microscopical examination. The writer prefers

^{*} Some analysts get better results with a mixture of ether and alcohol,

to have them slightly oleaginous, so that when applied to the slide no mountant need be used. In this case the crystals seem to stand out in wider contrast to the background than when cottonseed oil, the usual medium, is used.

If the crystals are, however, in a pulverulent condition, a drop of alcohol can be used as a mountant, or oil, as preferred. Mounted under a cover-glass they are examined under various powers of the microscope.

Figs. 272 and 273, Pl. XXXIX, show the typical appearance of pure lard stearin from a leaf lard of known purity, and Figs. 276, 277. and 278 illustrate beef stearin. These figures show distinctive crystallization of each form under the best conditions. The lard stearin crystals when thus obtained are flat rhomboidal plates cut off obliquely at one end, and are grouped irregularly, as if thrown carelessly together. The beef stearin crystals, on the other hand, are cylindrical rods or needles, often curved, with sharp ends, and are arranged as shown in fanshaped clusters. Conditions of crystallization are frequently such as not to show the sharp distinctions noted above. Both forms of crystals are at times apt to gather in clusters that at first sight appear somewhat similar, and are often misleading as to their true character. It is found almost invariably that the beef stearin crystals gather in clusters, radiating from a common center or point, often with a peculiar twisted appearance, breaking up into little fans. Lard crystals, it is true, do not always lie flat in irregular groups as shown in Fig. 272, but, as in Fig. 274, form clusters that, unless studied carefully, might at first sight be considered as identical with the fan shapes of the beef stearin already described. It will be seen, however, that if the best possible conditions are attained, the crystals of lard, instead of radiating from a point, are arranged more like feathers or alternate leaves on a branch, each crystal being given forth from another close at hand. Moreover, the lard crystals are themselves straight and not curved, the apparent curve in the appearance of the clusters being, on careful examination, especially under high power, seen to be chiefly due to several of these straight crystals arranged at angles to each other.

Even when the highest powers of the microscope are applied to the beef stearin crystals, they will always appear as cylindrical, sharp-pointed rods, some straight, others curved; while with the lard crystals they should be capable of showing the thin, flat, oblique-ended structure when examined with higher powers, even when they are arranged in the feathery clusters, the apparently pointed ends of some of the crystals

being due to the fact that the plates are viewed edgewise. This is apparent in Fig. 275, in which the crystals are magnified to 480 diameters.

According to Belfield, who was one of the earliest to employ the microscope for identification of foreign fat in lard, it is possible to detect well-defined crystals of both lard and beef stearin in mixtures crystallized out in the above manner from ether. Later investigators, however, find difficulty in getting both kinds of crystals in the final deposit, it being the more common experience that the character of the final crystals from a mixture of the two fats more often tends to one or the other forms of crystallization. Repeated crystallizations may change the character of the crystals and a number of such crytallizations should therefore be made before final judgment is passed.

The Iodine Number (p. 487).—This test is generally prefigured by the refractometer. Cottonseed oil will absorb about 108% of its weight of iodine, while beef fat will absorb about 37%.

ANALYSES OF SAMPLES ILLUSTRATING TYPES OF LARD, LARD SUBSTITUTES. AND MIXTURES.

				Butyro-refrac- tometer.		£		
	Nitric Acid Test.	Crystallization.	Bechi Reaction.	Temperature, Degrees.	Reading.	Variation from Standard.	Iodine Number.	Conclusion.
A	Slight color	Lard stearin	None	12.5	49.7	+0.1	58.1	Lard
A B	Red	"	***	42	50	+0.2	59.9	
Č	Slight color	46 16	**			+0.0		**
Ď	7, ,,	** **	66		50	+0.6		**
E	"	66 66	**	41.3		+0.8	64.6	"
D E F G	"	** **	**	42		+0.7	64.8	
Ğ	""	** **	**	42		-0.1	56.4	
H	Very slight color	Beef stearin	"			-3.8	37.3	
I	Deep-brown red	Few small bunches	Deep color	42	58.7	+8.9	108	Cottonseed oil
J	Red	Lard stearin	** **	43	50.5	+1.3	69.5	Lard and cotton- seed oil
K	Very slight color	Lard and beef stearin	None	43	48.5	-0.7	55-2	Lard and beef
L	Deep brown	Lard stearin	Deep color	43-5	51	+1.1	71.4	Lard and cotton- seed oil
M	Red	"	40 66	43.7	50.1	+1.3	66.7	
N	"	Lard and				+0.3		
		beef stearin					J	and cottonseed oil
			·	·				

Notes on the Above Table.—It will in general be noted that adulteration of lard with cottonseed oil alone is indicated by an abnormally high refractometer number, while the presence of tallow will result in an abnormally low refraction. But both adulterants may be present and a normal refraction result. In such a case the positive detection of one of them, such as the cottonseed oil by the Bechi or Halphen test, will-indirectly show the presence of the other (tallow), and this indirect proof will be confirmed by crystallization.

Samples A, B, and C give reactions corresponding to normal, pure lard. D, E, and F show somewhat high refractometer and iodine numbers, but give no direct reaction for cottonseed oil by the Bechi test. G, although showing low iodine and refractometer numbers, gives no evidence of the presence of tallow by crystallization. In fact, the crystals from this sample proved under all circumstances to be most clearly typical of pure lard, broad and flat plates with obliquely cut ends.

This sample was, in fact, pure leaf lard. It is generally true that a stiff, strictly pure leaf lard, which both by its consistency and by its low iodine and refractometer numbers might suggest the presence of beef fat, shows on crystallization much more definitely characteristic lard stearin than does a whole-hog lard, whose iodine and refractometer numbers are more nearly the normal standard.

In distinction from such leaf lard, a sample which may have a similar consistency and iodine and refractometer numbers, but which is composed of a whole-hog lard of a comparatively high iodine number, together with beef fat, gives unmistakable proof of its adulteration by its crystallization.

Effects of Feeding Hogs on Oil Cakes.—Fulmer,* Emmett and Grindley† and other investigators have found that feeding cottonseed meal to hogs causes the lard from these hogs to give a color with the Halphen test, but Tolman,‡ Farnsteiner§ and Polenske|| have shown that the lard does not contain phytosterol when examined by Bömer's phytosterol acetate method.

Lard from hogs fed on sesame cake has been shown to respond to the Baudouin test, but not to the phytosterol acetate test.

^{*} Jour. Am. Chem. Soc., 26, 1904, p. 837.

[†] Ibid., 27, 1905, p. 263.

[‡] Ibid., p. 589.

[§] Zeits. Unters. Nahr. Genuss., 11, 1906, p. 1.

^{||} Arb. Kaiserl. Gesundheitsamt., 22, 1905, p. 568.

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CHAPTER XIV.

SUGAR AND SACCHARINE PRODUCTS.

Nature and Classification of Sugars.—Of all classes of food materials the sugars from their great solubility are the most readily available, and on this account are very valuable as nutrients. As in the natural processes of digestion the starches and more difficultly digestible of the carbohydrates are converted into sugar and thus rendered assimilable, so by processes quite analogous to those that take place in the alimentary tract, the chemist converts these same carbohydrates into sugar as an end-point for purposes of definite determination.

The sugars are characterized by their sweet taste, their ready solubility in water, their power to rotate the plane of polarized light, and their insolubility in ether and absolute alcohol.

The sugars occurring commonly in food naturally divide themselves into two groups: First, the Saccharoses, or cane sugar group, having the composition $C_{12}H_{22}O_{11}$, of which the most prominent members are sucrose, maltose, and lactose; and, second, the Glucoses, or grape sugar group, expressed by the formula $C_6H_{12}O_6$, which includes dextrose, levulose and galactose, besides other less common sugars.

The members of both groups are intimately related. Thus by the ordinary process of so-called inversion sucrose, or cane sugar, belonging to group 1, is converted by the action of heat and dilute acid into two sugars, dextrose and levulose, which are members of group 2, in accordance with the following reaction:

$$\begin{array}{c} C_{12}H_{22}O_{11} + H_2O = C_6H_{12}O_6 + C_6H_{12}O_{6^*} \\ \text{Cane sugar} \end{array}$$
 Levulose

The same formula expresses also the result that takes place when lactose, or milk sugar, is heated with dilute acids, breaking up into dextrose and galactose.

Occurrence.—Sugars occur in roots, grasses, stems of plants, trunks of trees, leaves, and fruits, usually in the form of cane sugar, or sucrose, and of invert sugar (dextrose and levulose) mixed in varying proportions.

The following table from Buignet * shows the kind and amount of sugars occurring in some of the common fruits:

	Cane Sugar.	Reducing Sugar.	Acid.
Apricots	6.04	2.74	1.864
Pineapples	11.33	1.98	-547
English cherries	.00	10.00	.661
Lemons	.41	1.06	4.706
Figs	.00	11.55	-057
Strawberries		4.98	-550
Raspberries	2.01	5.22	1.380
Gooseberries	.00	6.40	1-574
Oranges		4.36	.448
Peaches (green)	.92	1.07	3.900
Pears (Madeleine)	.36	8.42	.115
Apples	5.28	8.72	1.148
"	2.19	5-45	.633
Prunes	5-24	2.43	1.288
Grapes (hothouse)	.∞	17.26	-345
" green	.∞	1.60	2.485

CANE SUGAR, OR SUCROSE.

Nature and Occurrence.—This, the most common of all the sugars, is nearly always understood by the unqualified term of sugar. It crystallizes in monoclinic prisms. Its specific gravity is 1.595. Its meltingpoint is about 160° C. Its specific rotary power $[\alpha]_D$ in solutions having a concentration of from 10 to 20 grams in 100 cc. is, according to Tollens, 66.48°. Sucrose is extremely soluble in water, which, when cold, will hold in solution twice its weight of the sugar.

Cane sugar is ordinarily derived from four sources—the sugar beet, the sugar cane, the maple tree, and the sorghum plant. The first two sources supply the principal output of commercial cane sugar, about two-thirds of the sugar on the market being furnished, according to Wiley, by the sugar beet and one-third by the sugar cane. It should be understood that the product sucrose, or cane sugar, is chemically the same whether derived from either of the above sources and thoroughly refined.

U. S. Standard Sugar is white sugar containing at least 99.5% of sucrose.

^{*} Ann. Chim. Phys., 59, 233.

The Sugar Cane (Saccharum officinarum) is cultivated principally in Louisiana and other southern states, in Cuba and the West Indies, and in the Hawaiian Islands. Its growth and cultivation form an industry in nearly all tropical countries.

Allen * has compiled the following table showing the composition of the juice of the sugar cane from different localities:

Locality and Kind of Cane.	Water.	Sugar.	Woody Fiber.	Salts.	Authority.
Martinique	72.1	18.0	9.9		Peligot
Guadaloupe	72.0	17.8	ý. 8	0.4	Dupuy
Havana	77.0	12.0	11.0		Casaseca
Cuba	65.9	17.7	16.4		Casaseca
Mauritius	69.0	20.0	10.0	1.0	Icery
Ribbon cane	76.73	13.39	9.07	-39	Avequin
Tahiti	76.08	14.28	8.87	•35	Avequin

The composition of raw cane sugar ash according to Monier is as follows:

RAW CANE SUGAR ASH.

Carbonate of calcium	. 49.00
" potassium	16.50
Sodium and potassium sulphate	. 16.00
Sodium chloride	. 9.00
Silica and alumina	9.50
	100.00

Manufacture of Cane Sugar.—The process of manufacturing raw sugar from sugar cane is briefly as follows: The juice is first extracted from the canes by crushing in roll mills and is freed from nitrogenous bodies, organic acids, etc., by the process of dejecation, which consists in heating to coagulate the albumin, and nearly neutralizing with milk of lime, the impurities being removed as a scum. The juice is then subjected to evaporation and crystallization, the raw, or muscovado sugar, which contains from 87 to 91 per cent of sucrose, being separated from the molasses, which is the mother liquor, by draining or by centrifugal.

Some of the best grade of muscovado, or raw sugar, is used as "brown sugar" without further refining, and much of the molasses is used as a table syrup and for cooking, while the lower grades of molasses are used in the manufacture of rum.

^{*} Com. Org. Analysis, 1, p. 315.

The following table from Thorpe * shows the average composition of raw and refined sugar:

	Cane Sugar.	Glucose ¹ .	Water.	Organic Matter.	Ash.
RAW SUGAR.					
Good centrifugal	96.5	0.75	1.50	0.85	0.40
Poor centrifugal	92.0	2.50	3.00	1.75	0.75
Good muscovado	91.0	2.25	5.00	1.10	0.65
Poor muscovado	82.0	7.00	6.00	3.50	1.50
Molasses sugar	85.0	3.00	5.00	5.00	2.00
Jaggary sugar	75.0	11.00	8.00	4.00	2.00
Manilla sugar	87.0	5-50	4.00	2.25	1.25
Beet sugar, 1st	95.0	0.00	2.00	1.75	1.25
Beet sugar, 2d	91.0	0.25	3.00	3-25	2.50
REFINED SUGAR.		1			
Granulated sugar	99.8	0.20	0.00	0.00	0.00
White coffee sugar	ό1.0	2.40	5.50	0.80	0.30
Yellow X C sugar.	87. 0	4.50	6.00	1.50	1.00
Yellow sugar	82.0	7.50	6.00	2.50	2.00
Barrel sugar.	40.0	25.00	20.00	10.00	5.00

¹ The term "glucose" includes sugars which reduce Fehling's solution, but are not necessarily optically active.

The following minimum and maximum figures are taken from analyses made by Babington † of twenty-two samples of brown sugar and thirty-one samples of molasses.

BROWN SUGAR.

Direct polarization. Invert "	84 - 27	to	87 - 29
Sucrose by Clerget	•	"	91.5
Reducing sugar	3	"	6
Moisture	-	"	6
Ash	0.8	"	3.0
MOLASSES.			
Direct polarization	30	to	50
Invert "	– 10	"	-21
Sucrose by Clerget	32	"	52
Reducing sugar	13	"	24
Moisture	29	"	32
Ash	0.5	"	4

^{*} Outlines of Industrial Chem., p. 383. † Can. Inl. Rev. Dept. Bul. 25.

The Sugar Beet (Beta vulgaris) is grown chiefly in France and Germany, and to a lesser extent in Holland and England. The successful growth of the sugar beet in the United States is confined mainly to California, Utah, and Nebraska, and the entire output of beet sugar in this country is comparatively small.

According to R. Hoffmann, sugar beets have about the following composition, three types being selected—first, those poor in sugar; second, those having a medium sugar content, and third, those rich in sugar:

COMPOSITION OF THE SUGAR BEET.

	Pirst Type.	Second Type.	Third Type
Water	89.20	83.20	75.20
Sugar	4.00	9.42 1.64	15.00
Nitrogenous compounds	1.00	1.64	2.20
Soluble	4.13	3-34	4.23
Insoluble (cellulose)	1.01	1.50	2.07
Ash	0.66	0.90	1.30
	100.00	100.00	100.00

The following is the mean composition of ten samples of California sugar beet:*

Per cent juice extracted	61.38
Specific gravity	1.062 to 1.075
Per cent of reducing sugar	0.91
Per cent of sucrose	14.38
Total solids calculated	16.58
Total solids weighed	
Per cent of ash	0.994

The composition of beet sugar ash according to Monier is as follows:

RAW BEET SUGAR ASH.

Carbonates of potassium and sodium	82.20
Carbonate of calcium	6.70
Potassium and sodium sulphate and sodium chloride	11.10
	100.00

Manufacture of Beet Sugar.—In making raw sugar from sugar beets the latter are first washed and sliced by machinery and the juice extracted

^{*} U. S. Dept. of Agric., Div. of Chem., Bul. 27, p. 202.

by diffusion or digestion with warm water. The juice is then clarified or defecated in much the same manner as that from the sugar cane, after which it is usually bleached with sulphur dioxide.

The subsequent evaporation and crystallization are carried out usually in vacuum pans, and the sugar separated out by centrifugals.

Beet sugar molasses is unfit for food, due to the presence of nitrogenous bodies, which give it a very unpleasant taste and smell.

Process of Refining.—In refining raw sugar, a syrup is made, which is subjected to centrifuging and further defecation, using lime, clay, liquid blood, calcium acid phosphate, and other substances as clarifiers. The juices are then filtered, first through cloth bags and then through bone char, after which they are evaporated and allowed to crystallize, the resulting granulated sugar being separated, as in the case of raw sugar, by centrifugal machines.

Granulated Sugar of commerce is without doubt the purest food product on the market, being generally 99.8% sucrose. It is usually treated with an extremely weak solution of ultramarine to counteract the natural yellow color.

The syrup from which the granulated sugar is separated forms the "golden," or "drip," syrup used on the table. Its typical composition is as follows: Sucrose, 40%; reducing sugars, 25%; water, 20%; organic matter, 10%; ash, 5%.

The dry sugars, whether white or brown, are rarely subjected to adulteration.

Maple Sap.—The sap of the maple tree, Acer saccharinum, or Acer barbatum, furnishes a sugar considerably prized for its peculiar flavor. The maple sugar industry is largely confined to the northeastern states and to Canada, and the maple sugar season is generally limited to six weeks or two months in the spring.

The following are minimum and maximum figures from the analyses of five samples of maple sap made in Massachusetts:

Specific gravity		
Sucrose	0.769 "	2-777
Reducing sugar	"	0.012

The ash of maple sap varies from 0.04 to 0.1 per cent. Albuminoids are present in amount varying from 0.008 to 0.03 per cent.

Maple Sugar and Syrup are made by simply boiling down the sap to the proper consistency, usually in open pans, and removing the scum with great care, since this contains nitrogenous matters that would cause fermentation in the finished product. Pure cane sugar is never commercially produced from the maple sap, since the refining process would deprive it of the flavor which gives to maple sugar the chief value.

McGill gives the following as the average analyses of six samples of maple syrup of known purity:

Saccharim-	Saccharim- eter Invert.		Ву С	opper.	Ash.		Solids.
			Reducing Sugar.	Cane Sugar.		Water.	
+62.2	-21.2	62.4	.42	63,36	•53	35 - 70	64.30

The variation in the composition of pure maple products is shown by the following table compiled by A. H. Bryan * from analyses published by Hortvet,† Jones,‡ and Winton§, and some sixty analyses made at the sugar laboratory of the Bureau of Chemistry, U. S. Department of Agriculture..

	Maple Sugar.			Maple Syrup.		
	Mini- mum.	Maxi- mum.	Average.	Mini- mum.	Maxi- mum.	Average
Water per cent	3.05	11.0		Not m	ore tha	n 32.00
Direct polarization	72.6	87.4		51.0	62.2	32000
Invert sugar "	1.16	8.37		0.34	9.17	
Lead number	1.83	2.48	2.23	1.19	2.03	1.49
Total ash per cent	0.64	1.32	0.91	0.46	1.01	0.66
Soluble ash "	0.33	0.67	0.46	0.21	0.63	0.38
Insoluble ash	0.20	0.87	0.46	0.14	0.56	0.23
Alkalinity of soluble ash		0.95	0.63	0.26	0.68	0.50
Alkalinity of insoluble ash		1.72	0.94	0.31	0.94	0.54
Ratio of insoluble to soluble ash	0.50	2.20	1.00	0.60	3.20	1.70
Iodine reaction			none	.		none
Polarization at 87° after inversion°V.		+2.0		- 240	+2.0	
Malic acid value	0.65	1.67	1.01	0.41	1.76	0.78

Partial ash analyses of maple products and brown sugar have been made by Jones || with the following maxima and minima results:

^{*} U. S. Dept. Agric., Bur. of Chem., Circular No. 40, p. 10.

[†] Jour. Am. Chem. Soc., 26, 1904, p. 1523.

Vt. Agric. Exp. Sta. Rep., 1904, p. 446; 1905, p. 315.

[§] Jour. Am. Chem. Soc., 28, 1906, p. 1204.

^{||} Loc. cit., 1905, p. 331.

		Number of Analysis.	100 Parts of Ash Contain			Ratio of			
			CaO.	K₂O.	SO ₃ .	CaO to KrO	CaO to SO ₂	K+O to SO	
Maple syrup:	Min Max	1 1	18.03	30.00 38.98	0.68	150	3.4	1.9	
Maple sugar:	Max Min Max	4	21.03	18.26	1.51	57	5.2	7.2 5.1	
Brown sugar:	Min Max		31.74 4.17 21.62	32.95 30.72 55.40	2.42 4.58 17.78	1 53 257 949	10.4 27 157	9-4 11 58	

^{*} Including one analysis by Hortvet.

U. S. Standards for Maple Products.—Maple Sugar is the solid product resulting from the evaporation of maple sap, and contains in the water-free substance not less than 0.65% of maple sugar ash.

Maple syrup is syrup made by the evaporation of maple sap or by the solution of maple concrete, and contains not more than 32% of water and not less than 0.45% of maple syrup ash.

Adulteration of Maple Sugar and Syrup.—The chief adulterants of maple sugar are brown, or molasses sugar, and white, or refined sugar, the latter being often used in mixture with burnt or inferior maple stock, which itself would be abnormally dark in color and of a rank taste. Maple syrup is commonly adulterated with a syrup made from refined cane sugar, less often with golden or drip syrup, or molasses. Glucose, which formerly was a common adulterant, is now seldom employed.

Refined Sugar or refined sugar syrup added to maple products, while not greatly affecting the polarization, diminishes the percentage of total ash and the lead number, as well as the malic acid value and ash constants. The ash of maple sugar should not be less than 0.64% and of maple syrup not less than 0.45%, while the lead number of maple sugar should not be less than 1.19.

According to analyses by Jones and Hortvet, brown sugar of various grades contains from 0.59 to 4.33% of total ash, some of the grades with low ash content, or syrups made from them, not being distinguishable from maple sugar or maple syrup respectively by this determination alone; the ratio of insoluble to soluble ash, however, is commonly higher in brown sugar than in maple products. It is frequently possible to identify brown, or molasses sugar, especially when it forms the larger portion of the alleged maple sugar or syrup, by the physical sense of taste. When the perfectly characteristic taste of brown, or molasses sugar, or of "drip syrup," so far predominates over the maple flavor as to be unmistakable,

especially in cases where the maple flavor is entirely lacking, one need have little hesitation in condemning the product.*

Glucose in maple products is detected by polarization both before and after inversion. A reading of the inverted solution much in excess of 3 degrees Ventzke at 87° C. furnishes evidence of the presence of this adulterant.

Sorghum (Andropogon sorghum, variety saccharatus) has for many years been grown quite extensively in the southern and western states, and used as a source of syrup, but only in recent years has it been found practicable to produce crystallized cane sugar from it on account of the presence of starch, uncrystallizable sugar, etc.

Much experimental work has been done of late along this line by the U. S. Department of Agriculture. The sorghum plant is as yet, however, a very small factor in the production of cane sugar, though much progress is being made.

The composition of the juice of the sorghum plant is shown by the following results of analyses of eleven varieties made by Hardin:

Total solids	15-97	to	18.71
Specific gravity	1.0656	"	1.0775
Solids not sugar	5.02	"	10.63
Cane sugar	2.81	"	8.01
Reducing sugars	3.87	"	7-55

Some varieties of sorghum juice have been known to contain 15 or even 17 per cent of sucrose.

In making syrup from sorghum, the ripe canes are crushed, the juice is heated with milk of lime, and the scum removed. The juice is then concentrated usually in open pans to the required consistency.

GRAPE SUGAR, OR DEXTROSE.

Dextrose $(C_6H_{12}O_6+H_2O)$, designated d-glucose by Fischer and known in its commercial form as starch sugar, occurs in honey with levulose, and in fruits with both levulose and cane sugar. It is produced by the action of dilute acids or of certain ferments on starch, dextrin, or cane sugar. Grapes contain about 15% of dextrose. Anhydrous

^{*} The sense of taste, if properly cultivated, and with its limitations recognized, should be entitled to as much consideration as the other senses in forming an opinion. Taste and smell are often very useful factors in detecting adulterants, but should of course be used with discretion.

[†] U. S. Dept. of Agric., Div. of Chem., Bul. 37, p. 75.

dextrose is soluble in 1.2 parts of cold water. It is soluble in alcohol, but less so than cane sugar. It is much less sweet than cane sugar.

The specific rotary power of dextrose is

$$[\alpha]_D = 52.3, \quad [\alpha]_j = 58.$$

A normal solution of dextrose on the Soleil-Ventzke scale polarizes at 78.6°. For the commercial preparation of dextrose see p. 576.

U. S. Standards for Various Sugars.—Standard 70 sugar, or brewers' sugar, is hydrous starch sugar containing not less than 70% of dextrose, and not more than 0.8% of ash.

Standard 80 sugar, climax, or acme sugar, is hydrous starch sugar containing not less than 80% of dextrose, and not more than 1.5% of ash.

Standard anhydrous starch sugar is anhydrous starch sugar containing not less than 95% of dextrose without water of crystallization, and not more than 0.8% of ash.

The ash of these standard products consists almost entirely of chlorides and sulphates of lime and soda.

LEVULOSE.

Levulose, also known as d-fructose and l- δ -fructose, occurs in foods as the product of inversion of cane sugar. It is prepared by the action of dilute acids on inulin. Normally it is in the form of a syrup, but with extreme care pure anhydrous levulose can be obtained. Diabetene is a commercial form of dry levulose. Levulose is formed with dextrose in the inversion of cane sugar (p. 565), and with dextrose occurs in honey and in many fruits. The specific rotary power of levulose varies with the temperature. At 15° C. $[\alpha]_D = -98.8^\circ$, decreasing by 0.6385° for each degree increase in temperature. Its left-handed reading on the Ventzke sugar scale at 15° C. is equivalent to 148.6°. Levulose is sweeter than dextrose. Its reducing power on Fehling's solution is assumed to be the same as that of dextrose.

MALT SUGAR, OR MALTOSE.

Maltose (C₁₂H₂₂O₁₁+H₂O) is of little importance from the standpoint of the food analyst, excepting as an ingredient of commercial glucose, and as being the sugar produced by the action of ptyaline, the ferment of the saliva on the starch of food in the ordinary process of digestion. When gelatinized starch is subjected to treatment with malt extract at 55° to 60° C., it is converted into dextrin and maltose as follows:

$$\begin{array}{c} \mathbf{C_{18}H_{30}O_{15} + H_{2}O = C_{6}H_{10}O_{5} + C_{12}H_{22}O_{11}.} \\ \mathbf{Starch} & \mathbf{Dextrin} & \mathbf{Maltose} \end{array}$$

In its commercial preparation maltose is separated from dextrin by crystallization in alcohol. By the action of weak acids and heat both dextrin and maltose are further converted into dextrose.

Maltose usually crystallizes in minute needles, and its molecule of water is expelled at 110° C. It is somewhat less soluble in water than dextrose. It is slightly soluble in alcohol, though less than sucrose. Solutions of maltose possess the property of birotation; i.e., when freshly prepared they do not at once assume their true optical activity. The rotation of a freshly prepared solution of maltose increases on standing, requiring several hours to reach its maximum. The specific rotary power, according to O'Sullivan, of anhydrous maltose is $[\alpha]_D = 139.2$, $[\alpha]_i = 154.5$. For hydrated maltose $[\alpha]_D$ would thus be 132.2.

A normal solution of maltose on the Soleil-Ventzke scale should polarize at 198.8°.

DEXTRIN. COMMERCIAL GLUCOSE.

DEXTRIN, $(C_0H_{10}O_5)_m$, possesses more the nature of a gum than of a sugar, and is sometimes called British gum. It is said to occur naturally in the sap of various plants, but this is not definitely assured.

It undoubtedly occurs in beer and in bread crust, and is one of the constituents of commercial glucose. Like starch, it is convertible by hydrolysis with acid into dextrose. By treatment of starch with malt extract or diastase, starch is converted into dextrin and maltose, these two bodies being separated, in the commercial preparations of dextrin, by repeated treatment with alcohol.

Dextrin is an uncrystallized, colorless, tasteless body, capable of being pulverized. It is readily soluble in water, slightly soluble in dilute alcohol, but insoluble in alcohol of 60% or stronger. It is not colored by iodine, and exercises no reducing action on alkaline copper solution. Its specific rotary power is $[\alpha]_D = 200$, $[\alpha]_{j=202}$.

Amylodextrin, erythrodextrin and achroodextrin are intermediate products formed in the transformation of starch into dextrose. Amylodextrin is colored purple and erythrodextrin red by iodine solution, while achroodextrin produces no coloration. It is probable that some of these dextrins are not simple substances.

Commercial Glucose, otherwise known as mixing syrup, crystal syrup, and starch, or corn syrup, is a heavy, mildly sweet, colorless, semi-fluid substance, having a gravity of 40° to 45° Baumé. It is largely used as an adulterant of maple syrup, molasses, honey, drip syrup, and jellies and jams, and as an ingredient of confectionery.

In France and Germany it is made from potato starch, but in the United

States mainly from corn starch. The conversion is effected by boiling with dilute sulphuric or hydrochloric acid, after which the acid is neutralized with marble dust, or sodium carbonate respectively, the juice is filtered through bone black, and finally concentrated by evaporation, the degree of conversion and of concentration depending on whether the liquid glucose or the solid dextrose is wanted for the final product. The end product obtained by complete conversion is the dry commercial grape sugar, or dextrose, which is purified by repeated crystallization.

Commercial glucose is a mixture of dextrin, maltose, and dextrose of the following varying composition:

Dextrin	29.8%	to 45.3%
Maltose	4.6%	" 19.3%
Dextrose	34-3%	" 36.5%
Ash	0.32%	" 0.52%
Water	14.2%	" 17.2%

Calcium sulphate is usually found in the ash if sulphuric acid was used for conversion.

Solid commercial grape sugar, or dextrose, has the following composition:

Dextrin	o%	9.1%
Maltose	o%	1.8%
Dextrose	72%	99.4%
Ash	0.3%	0.75%
Water	$\circ.6\%$	17.5%

U. S. Standard glucose, mixing glucose, or conjectioners' glucose, is colorless glucose, varying in density between 41° and 45° Baumé, at a temperature of 100° F. (37.7° C.). It conforms in density, within these limits, to the degree Baumé it is claimed to show, and for a density of 41° Baumé contains not more than 21% of water, and for a density of 45° not more than 14%. It contains on a basis of 41° Baumé not more than 1% of ash, consisting chiefly of chlorides and sulphates of lime and soda.

Healthfulness of Glucose.—The analyst alleging commercial glucose as an adulterant is frequently asked in court as to its healthfulness, so that the following conclusions of a committee appointed some years ago by the National Academy of Sciences to ascertain among other things whether there is any danger attending the use of this product in food are in point: "First, that the manufacture of sugar from starch is a long-established industry, scientifically valuable and commercially important;

second, that the processes which it employs at the present time are unobjectionable in their character and leave the product uncontaminated; third, that the starch sugar thus made and sent into commerce is of exceptional purity and uniformity of composition and contains no injurious substances; and fourth, that though having at best only about two-thirds the sweetening power of cane sugar, yet starch sugar is in no way inferior in healthfulness, there being no evidence before the committee that maize starch sugar, either in its normal condition or fermented, has any deleterious effect upon the system, even when taken in large quantities."

MILK SUGAR, OR LACTOSE.

Lactose (C₁₂H₂₂O₁₁+H₂O) is prepared commercially from skimmilk by coagulating with rennet and digesting the whey with chalk and aluminum hydroxide. The insoluble matter is filtered out, and the filtrate is concentrated *in vacuo* to a syrup, which, on standing, yields crystals of lactose. The product is purified by repeated crystallization.

Lactose ordinarily crystallizes in rhombic, hemihedral crystals. Its specific gravity is 1.525. Its water of crystallization is lost by drying at 130° C. It is soluble in 6 parts of cold water, and in 2½ or less of boiling water. It is insoluble in absolute alcohol and ether. It has a very slightly sweet taste.

The specific rotary power of milk sugar, after remaining in solution long enough to overcome its birotation, is

$$[\alpha]_{0} = 52.5.$$

In the ordinary souring of milk the lactose becomes converted into lactic acid.

On heating lactose with dilute acids it undergoes inversion, forming dextrose and galactose in accordance with the formula given on p. 565, illustrating the inversion of cane sugar.

Milk sugar is of considerable importance by reason of the large amount used of late in the preparation of modified milk for infant feeding.

Grape sugar and cane sugar are to be looked for as adulterants of milk sugar.

The purity of milk sugar is best established by titrating against Fehling's solution, 10 cc. of which are equivalent to 0.067 gram of lactose.

RAFFINOSE.

Raffinose, C₁₈H₃₂O₁₆5H₂O, is a sugar belonging neither to the saccharose nor the glucose group, but to the so-called saccharoid group, the other members of which do not occur in foods.

Raffinose occurs in beet root molasses to the extent of from 3 to 4 per cent. It is a crystalline, slightly sweet substance, soluble in water and slightly soluble in alcohol. It does not reduce Fehling's solution, but readily undergoes fermentation. On inversion it splits up into levulose and melibiose (C₁₂H₂₂O₁₁).

The melting-point of raffinose is 118° to 119° C. Its specific rotary power $[\alpha]_D = +104.5$ at a temperature of 20° C.

THE POLARISCOPE AND SACCHARIMETRY.

A full discussion of the principles of polarized light and even a detailed description of their application to the polariscope will not be given here, but the reader who wishes full information along this line is referred to the various text-books, and especially to those of Tucker, Spencer, and Landolt,* in which various forms of polariscopes are described and their underlying principles discussed.

The Soleil-Ventzke Saccharimeter is the one most commonly used in this country, being adopted as the standard for all United States government work. Fig. 102 shows this instrument, known as the half-shadow

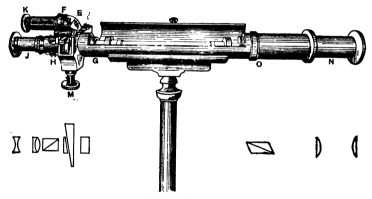


Fig. 102.—Single-wedge Saccharimeter.

apparatus, in its simplest form with a single movable wedge in its compensating system.

An excellent light for work with this instrument is that furnished by the Welsbach burner, a convenient form of lamp being shown in Fig. 111, in which the burner is inclosed in a sheet-metal chimney of suitable construction. An argand, gas, or kerosene burner may however be used,

^{*} See references, p. 651.

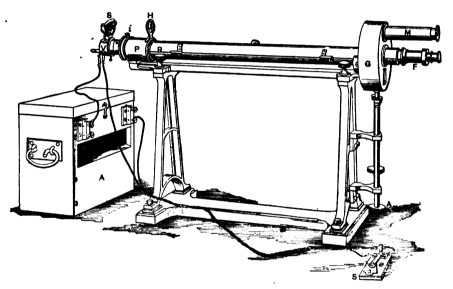
and in a late form of Schmidt and Haensch instrument, Fig. 103, a specially constructed incandescent electric lamp is supplied.

The Single-wedge Saccharimeter.—The following description of the saccharimeter and directions for its use are from the revised regulations of the U. S. Internal Revenue Department. The tube N, Fig. 102, contains the illuminating system of lenses and is placed next to the lamp; the polarizing prism is at O and the analyzing prism at H. The quartz wedge compensating system is contained in the portions of the tube marked FEG and is controlled by the milled head M. The tube J carries a small telescope, through which the field of the instrument is viewed, and just above is the reading-tube K, which is provided with a mirror and magnifying lens for reading the scale.

The tube containing the sugar solution is shown in position in the trough between the two ends of the instrument. In using the instrument the lamp is placed at a distance of at least 200 mm. from the polarizing end; the observer seats himself at the opposite end in such a manner as to bring his eye in line with the tube J. The telescope is moved in or out until the proper focus is secured to give a clearly defined image. when the field of the instrument will appear as a round, luminous disk, divided into halves by a vertical line passing through its center. and darker on one half of the disk than on the other, when the compensating quartz wedge is displaced from the neutral position. If the observer, still looking through the telescope, will now grasp the milled head M and rotate it first one way and then the other, he will find that the appearance of the field changes, and at a certain point the dark half becomes light and the light half dark. By rotating the milled head delicately backward and forward over this point he will be able to find the exact position of the quartz wedge operated by it, in which the field is neutral, or of the same intensity of light on both halves. The three different appearances presented by the field are shown in Fig. 106, opposite page 582,

One of the compensating quartz wedges is fixed and the other is movable, sliding one way or the other according as the milled head is turned, so that for different relative positions of the two wedges a different thickness of quartz is interposed in the path of the polarized ray. By this means the amount of the rotation which the sugar solution or other optically active substance examined exerts upon the light polarized by the prism at O may be, as it were, counteracted by varying the relative position of the wedges.

With the milled head set at the point which gives the appearance of the middle disk shown in Fig. 106, the eye of the observer is raised to the reading tube K, which is adjusted to secure a plain reading of the divisions, and the position of the scale is noted. It will be seen that the scale proper is attached to the quartz wedge, which is moved by the milled head; and attached to the other quartz wedge is a small scale called a vernier, which is fixed, and which serves for the exact determination of the position of the movable scale with reference to it. On each side of the zero line of the vernier a space corresponding to nine divisions of the movable scale is divided into ten equal parts. By this device the fractional part of a degree indicated by the position of the zero line is ascertained in



Fro. 103.—Single-wedge Soleil-Ventzke Saccharimeter, mounted on Bock Stand and provided with Incandescent Electric Lamp.

tenths; it is only necessary to count from zero until a line is found which makes a continuous line with one on the movable scale.

With the neutral field, as indicated above, the zero of the movable scale should correspond closely with the zero of the vernier, unless the zero point is out of adjustment.

Adjusting the Instrument.—If the observer desires to secure an exact adjustment of the zero of the scale, or in any case if the latter deviates more than one-half of a degree, the zero lines are made to coincide by moving the milled head and securing a neutral field at this point by

means of the small key which comes with the instrument, and which fits a small nipple on the *left*-hand side of *F*, the fixed quartz wedge of the compensating system. This nipple must not be confounded with a similar nipple on the *right*-hand side of the analyzing prism *H*, which it fits as well, but which must never be touched, as the adjustment of the instrument would be seriously disturbed by moving it. With the key on the proper nipple it is turned one way or the other until the field is neutral. Unless the deviation of the zero be greater than 0.5° it will not be necessary to use the key, but only to note the amount of the deviation, and for this purpose the observer must not be content with a single setting, but must perform the operation five or six times and take the mean of these different readings. If one or more of the readings show a deviation of more than 0.2° from the general average they should be rejected as incorrect. Between each observation the eye should be allowed a moment of rest.

The Scale usually has 110 equal divisions on one side of the zero for reading right-handed polarization, and 20 equal divisions on the other side for left-handed polarization. The scale is an arbitrary one, based on the plan that a normal aqueous solution of pure cane sugar (26.048 grams made up to 100 cc.) will read exactly 100° or divisions to the right of the zero.

The accuracy of various portions of the scale may be verified by quartz control plates of varying thickness, usually mounted in tubes, the correct polariscopic reading of each of which plates has been accurately determined, this reading being as a rule marked on the tube. As the sugar value of such a quartz plate varies with the temperature, the temperature at which the particular reading marked thereon applies is usually specified, and in many cases a table giving its exact value at different temperatures from 10° to 35° accompanies the plate.

The Double-wedge Saccharimeter is shown in Fig. 104, the arrangement of the optical parts being also shown.

In this instrument the two sets of wedges employed are of opposite optical properties, so that extreme accuracy may be arrived at by making the readings with both, the inaccuracies of one being compensated for by the other. Ordinarily in using this form, one movable wedge, say the one controlled by the right-hand milled screw head, is set at zero, while the reading of the sugar solution or other substance to be polarized is made with the other movable wedge.

The Trible-shadow Saccharimeter.—The latest form of saccharimeter

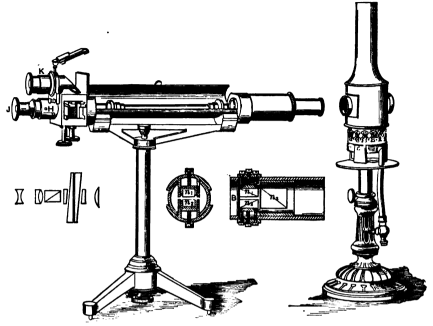


Fig. 104.—Double-wedge, Triple-shade Soleil-Ventzke Saccharimeter.

is the triple-shadow instrument, the construction of the polarizer being shown in Fig. 105.

I

In this form the analyzer is the same as in the foregoing instruments, but the polarizer consists of one large and two small Nicol prisms I, II, and III, the construction and arrangement being such that when the compensating wedges are at the neutral point, sections 1, 2, and 3 of the circular field (corresponding respectively to the prisms I, II, and III) are evenly lighted, forming a circular uniformly colored field, while in any other position of the wedges section 1 is dark while 2 and 3 are light or vice versa. The accompanying diagram, Fig. 106, shows the appearance of the field of this instrument in the three positions of the quartz wedge, viz., at the neutral point and at both sides thereof.

The lamp used for illumination should be separated from the polariscope on account of the influence of its heat on the readings. This is best accomplished by having the lamp in a separate compartment from the polariscope, so

that both are on opposite sides of a partition, an opening in which transmits the light. In any event some kind of screen should be interposed between the two. Best results are obtained if the room in which the observations are made is dark.

Comparisons of Scales of Various Polariscopes.—Besides the Soleil-Ventzke instrument, there are various other forms of polariscope. Among the best known of these are Laurent's, Wild's, and Duboscq's, all of which are made with scales reading in circular degrees, while in some cases modified forms have scales in which, like the Soleil-Ventzke, percentages of sugar are directly read off. Some instruments are provided with double scales reading both circular degrees and percentages of sugar, and in certain of the Duboscq instruments additional scales for percentages of milk sugar and diabetic sugar are provided.

In the Wild, Duboscq, and Laurent instruments the source of light is the sodium flame, yielding what is termed a monochromatic light. This is produced by fused sodium chloride passing through a Bunsen flame, various mechanical devices being employed for making the light continuous. In the Ventzke instrument, as was stated above, the ordinary light from a bright gas or oil flame is used.

For convenience in conversion of readings on one instrument to their equivalents on other scales, the following factors can be used:

```
=0.3468° angular rotation D.
=2.8835° Ventzke.
1º Ventzke
1° angular rotation D
                           =2.6048° Wild (sugar scale).
1º Ventzke
                           =0.3840° Ventzke.
1º Wild (sugar scale)
                           =0.1331° angular rotation D.
                           =0.7511° Wild (sugar scale)
1° angular rotation D
1º Laurent (sugar scale) = 0.2167° angular rotation D.
1º angular rotation D = 4.6154° Laurent (sugar scale).
                           =0.2167° angular rotation D.
1º Soleil-Duboscq
                           =0.2450°
10 "
             "
                                       Soleil-Ventzke.
                            =0.620°
   "
             "
                             =1.619°
                                       Wild.
                             = 1.608°
1° Soleil-Ventzke
                                       Soleil-Duboscq (old scale).
                           -1.593°
                                                       (new scale).
                           =0.6ii°
ro Wild
                                                       (Wild normal weight 10).
                           =1.223°
                                                                              20).
```

Normal Weights of Sugar for Different Instruments.—The following normal weights (number of grams in 100 cc. at 17.5° C.) are those on which the scales of the various instruments are based: Soleil-Ventzke, 26.048; Soleil-Duboscq 16.35 (formerly 16.19); Wild, usually, 10 or 20; Laurent, 16.19.

The International Commission for Uniform Methods in Sugar Analysis has decided to use for the Ventzke scale 26 grams and make up at 20° C. to 100 metric cc., which figures are approximately equivalent to 26.048 grams made up to 100 Mohr cc.

Specific Rotary Power.—This is a theoretical term to express a standard by which the various optically active substances may be compared, and is understood to mean the amount in angular degrees through which the plane of polarization of a ray of light of stated wave length is rotated by I gram of a given substance in aqueous solution of I cc. and forming a column I decimeter in length. The actual rotary power of a solution varies directly with the length of the column traversed by the light, with the concentration of the solution, and with the wave length of light, hence the need of a purely theoretical basis for purposes of comparison.

The specific rotary power is usually expressed as $[\alpha]_D$ or $[\alpha]_j$, the letters D or j indicating the character of the light. Thus, D indicates the monochromatic light obtained from the sodium flame, named from the D line of Fraunhofer in the yellow portion of the spectrum, while j (from the French *jaune*) indicates what is known as the *transition tint*, the rose-purple color produced when ordinary white light passes through the polarizer and analyzer, placed with their principal sections parallel to each other and with a plate of quartz 3.75 mm. thick interposed between them.*

The specific rotary power is determined as follows:

$$[\alpha]_D$$
 or $[\alpha]_i = \frac{100a}{cl}$,

where a = observed angular rotation,

c = grams of the substance in 100 cc. of the solution, and

l=length of the observation-tube in decimeters; or, in cases where, instead of the grams per 100 cc., the percentage composition is known (expressed by p=grams of the substance in 100 grams of the solvent),

and the specific gravity (expressed by d), then $[\alpha]_D$ or $[\alpha]_i = \frac{100a}{pdl}$.

Birotation.—In polarizing solutions of all the common sugars other than sucrose the phenomenon of birotation should be taken into account, whereby a change in optical activity is shown by standing. Thus, solutions of dextrose, levulose, and lactose polarize much higher when freshly prepared than after long standing, requiring in some instances several hours before the lowest or normal figure is reached. Maltose, on the other hand, increases in polarization after standing in solution. By

^{*}Some confusion is caused by the adoption of the characters D and j, since both would naturally seem to indicate yellow light. The so-called transition tint above defined is, however, complementary to the mean yellow, or jaune moyen, and it is the complementary color and not the yellow itself that is indicated by the character j

Fig. 106.—Appearance of the Field in the Half-shade (above) and Triple-shade (below) Saccharimeter.



boiling the solution it may be at once brought to its correct reading. The desired result may also be accomplished by adding a few drops of ammonia, either treatment being resorted to before the solution is made up to the required volume.

ANALYSIS OF CANE SUGAR AND ITS PRODUCTS.

Qualitative Tests for Sucrose.—(a) Polariscope Test.—The substance to be tested, if not already in solution, is dissolved in water, and if the solution is not perfectly clear, is clarified by the addition of alumina cream or by subacetate of lead (p. 586) and filtered. An observation tube is filled with the clear solution and the polariscope reading noted. A measured portion of the same solution is then treated with one-tenth its volume of concentrated hydrochloric acid and is subjected to inversion (p. 588), after which the same tube as before is filled with the inverted solution and a second reading obtained, one-tenth of the observed reading being added for the true invert polariscopic reading. If the two readings are virtually the same, sucrose is absent, but, in the presence of sucrose, the second reading will be considerably lower than the first or may even be to the left of the zero.

(b) Test with Nitrate of Cobalt.*—Prepare a 5% solution of cobaltous nitrate, and a 50% solution of potassium hydroxide. If the sugar solution to be tested contains dextrin or gums, these should first be removed by treatment with alcohol. 15 cc. of the sugar solution to be tested are mixed with 5 cc. of the cobaltous nitrate reagent, and 2 cc. of the potassium hydroxide solution are added. Sucrose produces under these conditions a permanent amethyst-blue color, while dextrose gives at first a turquoise-blue passing over into light green. In a mixture of the two sugars the color due to sucrose will predominate.

According to Wiley, I part of sucrose in 9 parts of dextrose may be detected by this test.

ANALYSIS OF CANE SUGAR.—In the case of commercial granulated or loaf sugar the sucrose determination is usually all that is necessary to determine its purity, and the same is true, as a rule, of the powdered white sugars. A fairly complete analysis of raw or brown sugar consists in the determinations of moisture, sucrose, invert sugar, ash, organic non-sugars, and quotient of purity. Care should be taken that the portion subjected to analysis is a fair representation of the whole, and is perfectly homogeneous.

Determination of Moisture.—2 to 5 grams of the sample are dried in a flat, tared metal dish, to constant weight *in vacuo*, or in a McGill oven* in a current of air, at about 70° C., at which temperature levulose is not decomposed. For ordinary purposes sufficiently accurate results may be obtained by the A. O. A. C. method of drying to constant weight at 100° C. in a water oven.

Determination of the Ash.—The residue from the moisture determination is burned slowly and cautiously over a low flame until frothing has ceased. Afterwards increase the flame and ignite to a white ash at a low, red heat.

In igniting saccharine substances which contain an appreciable amount of cane sugar, the contents of the dish will swell up and froth, unless great care be taken, to such an extent as to flow over the sides of the dish, occasioning loss and inconvenience. Such frothing may be largely held in check by directing the flame at first down from above upon the pasty mass, instead of from under the dish as ordinarily, till all is reduced to a dry char, afterwards continuing the ignition from below in the usual manner.

Organic Non-sugars.—These consist mainly of compounds of organic acids, together with gum, coloring matter, albuminous bodies, etc. They are determined by difference between 100% and the sum of the sucrose, invert sugar, moisture, and ash.

Quotient of Purity.—By this term is meant the percentage of pure sugar in the dry substance. It is calculated by dividing the per cent of sucrose by the percentage of total solids and multiplying the result by 100.

Determination of Sucrose by the Polariscope.—Reagents.—Subacetate of Lead Solution.†—This is prepared by boiling for half an hour 430° grams of normal lead acetate, 130 grams of litharge, and 1000 cc. of

^{*} A. McGill, Laboratory of Inland Revenue, Ottawa, Canada, has devised a forced-draft water-oven for drying at temperatures between 60° and 90° C. The oven is heated by means of ordinary gas-burners, and the temperature is controlled by introducing at the bottom of the oven a blast of air from a blower run by a small water-motor. Before discharging into the oven, the air-tube enters the water-chamber and is coiled a number of times in order to sufficiently warm the air before it enters the oven. The exit end of the air-tube is covered with a concavo-convex disk in order to distribute the blast and to prevent harmful currents. By regulating the burners and the flow of air, a fairly constant temperature can be obtained. The bottom of the oven is curved instead of flat, to prevent bumping when the water is boiling; a perforated plate serves as a false bottom.

[†] U. S. P. lead subacetate may be used. This is sometimes sold under the name of Goulard's extract.

water. The mixture is allowed to cool and settle, when the supernatant liquid is diluted to 1.25 specific gravity with recently boiled water.

Alumina Cream is prepared by dividing a cold, saturated, aqueous solution of alum into two unequal portions, to the larger of which add a slight excess of ammonia. Then add by degrees the remaining portion to a faint acid reaction.

Process.—If the Soleil-Ventzke polariscope is to be used, weigh out 26.048 grams of the sugar, which may conveniently be done in the German-silver, tared tray especially designed for this purpose, and which accompanies the Schmidt and Haensch polariscope, Fig. 107. If any other instrument than

FIG. 107.—German-silver Sugar-tray with Tare.

the Soleil-Ventzke is employed, weigh out the standard or normal weight for that instrument (see p. 583). Transfer the sugar by washing to a 100-cc. graduated sugar-flask, and if the solution is perfectly clear, as would be the case with a refined sugar, make up to the mark and shake to insure a uniform solution. If the solution is slightly turbid, or more or less opaque or dark-colored, a clarifier must be added before making up to the mark to obtain a clear solution for polarization. The kind

Fig. 108.—A Convenient Sugar-scale.

and amount of clarifier to be used depends on the nature of the sugar solution, and experience will soon indicate what is best adapted to given conditions. If the turbidity is only slight, from 5 to 10 cc. of alumina cream alone will often prove sufficient. In case of a very opaque solution, 10 cc. of subacetate of lead solution will nearly always suffice.

For additional details as to clarification see page 614. under Molasses. After adding the clarifier, the flask is filled to the mark with water and shaken, the solution being poured upon a dry filter and the first few cubic centimeters of the filtrate rejected. A 200-mm. observation-tube is filled with the clear sugar solution and the polarization noted. If sucrose is the only optically active substance present, the direct reading on the polariscope will indicate its percentage.

Process of Inversion.—In the presence of invert or other sugars the normal solution as above prepared is subjected to inversion as follows: Free a portion of the solution from lead by treating with anhydrous sodium carbonate, sodium sulphate or potassium oxalate, filter, place 50 cc. in a 100-cc. flask, add 25 cc. of water and little by little, while rotating the flask, 5 cc. of 38.8% hydrochloric acid. Heat in a water bath at 70° C., so that the solution in the flask reaches 67° to 69° C. in two and one-half to three minutes. Maintain at 69° C. during seven to seven and one-half minutes, making a total time of heating of ten minutes. Remove the flask, cool the contents rapidly to 20° C., and dilute to 100 cc. Polarize this solution in a 200-mm. tube provided with a lateral branch and a water jacket, passing a current of water around the tube to maintain a temperature of 20° C.

The inversion may also be accomplished by allowing a mixture of 50 cc. of the clarified solution, freed from lead, and 5 cc. of the acid to stand for 24 hours at not less than 20° C. or for 10 hours at not less than 25⁶.

The sucrose is obtained by the following formula of Clerget, based on the rotation of cane sugar before and after inversion,

$$S = \frac{100(a-b)}{142.66-t/2},$$

where S=per cent of sucrose, a=direct polarization, b=invert polarization, and t=temperature. Note that if the direct polarization is to the right or positive, and the invert to the left or negative, then a-b would be the sum of the two polarizations.

In many cases where it is almost impossible to obtain a colorless solution for polarization in the 200-mm. tube, a 100-mm. tube may be employed, and the readings multiplied by 2, or half the normal weight,* viz., 13.024 grams, of the sample may be taken and made up to 100 cc., the 200-mm. tube employed, and the readings multiplied by 2.

^{*}Wherever the term "normal weight" occurs hereafter will be meant, unless otherwise noted, the normal weight of sugar for the Soleil-Ventzke polariscope, viz., 26 grams, and by a "normal solution" will be meant 26 grams in 100 cc. of water. Clerget's formula, as originally worked out by him, was not based on this normal weight, but on 16.35 grams. It is, however, applicable to 26 grams.

The determination of sucrose by the Clerget formula is applicable to all mixtures of the common sugars excepting those in which lactose, or milk sugar, is present.

Theory of Inversion.—On p. 565 a reaction is given showing that when sucrose is subjected to inversion by the action of dilute acids or of invertase or yeast it splits up into the two sugars dextrose and levulose. forming equal quantities of each. The dextrose is, however, dextrorotary and the levulose lævorotary. Invert sugar is the term applied to the mixture of dextrose and levulose formed by the inversion of sucrose. The specific rotary power of sucrose varies so little with the temperature as to be regarded for practical purposes as constant. At 87° a solution of invert sugar polarizes at zero. This is due to the fact that the rotary power of levulose, unlike that of sucrose and dextrose, varies with the temperature. At from 87° to 88° the left-handed rotation of the levulose balances the right-handed rotation of the dextrose in the invert sugar, hence the zero reading. As the temperature decreases from 87°, the rotary power of the levulose proportionally increases, till at oo the normal invert sugar solution would polarize 44° to the left of the zero. On these facts Clerget's formula (p. 583) is based, assuming that a normal solution of pure cane sugar polarizes at +100°, while after inversion the reading for o° temperature would be -44° and would decrease half a degree for each degree in temperature above o°. Thus at 20° the invert reading, would be -34.

Detection of Invert Sugar.—Methyl-blue Test.—This test depends on the decolorization of methyl blue by invert sugar. 20 grams of sugar are dissolved in water and made up to 100 cc. If the solution is not clear, sufficient subacetate of lead solution is added to clarify before making up to the mark, and the solution is filtered. Add to the filtrate enough 10% sodium carbonate solution to make alkaline, and filter a second time. Take about 50 cc. of the filtrate in a casserole, add 2 drops of a 1% solution of methyl blue, and boil over a free flame, noticing particularly the time the solution begins to boil.

If the color disappears in one minute after boiling, there is present at least 0.01% of invert sugar. If it is not completely decolorized by three minutes' boiling, no invert sugar is present.

Determination of Invert Sugar in Cane Sugar Products by the Polariscope.—While invert sugar is best determined by Fehling's solution as described elsewhere, it may be approximately estimated by the polari-

scope, though less satisfactorily. On p. 626 a method is given for the determination of levulose by polariscopic readings at two different temperatures. Since invert sugar is composed of equal parts by weight of dextrose and levulose, the percentage of levelose multiplied by 2 would give that of invert sugar.

Test for Ultramarine in Sugar.*—A large amount of the sugar is dissolved in water and the coloring matter is allowed to settle out, washing the residue several times by decantation. On treatment with hydrochloric acid, the blue color is discharged if due to ultramarine.

SUGAR DETERMINATION BY COPPER REDUCTION.

Various convenient methods of determining sugars depend on the readiness with which certain of them, known as reducing sugars, act on copper salts, especially on the tartrate of copper, reducing it to cuprous oxide.

This reducing power is exercised in a definite degree under fixed conditions, so that the amount of reducing sugar present may be accurately determined. Of the common sugars, sucrose is the only one that has no direct reducing action, but on undergoing inversion it is converted into reducing sugars, which are readily determined.

Use of Fehling's Solution.—There are various well-known mixtures of copper sulphate, tartaric acid salts (usually Rochelle salts or cream of tartar), and alkalies, called after chemists who have employed them in the determination of the reducing sugars, each one possessing certain advantages, but none have become so widely adopted as Fehling's solution, the use of which in one form or another is now well-nigh universal.

There are a number of methods by which Fehling's solution is employed for this purpose, both volumetric and gravimetric. The former are simpler and quicker of manipulation, and thus are preferable for commercial work where extreme accuracy is not required. The gravimetric methods are usually considered more delicate and accurate, calling for less skill, but more time in arriving at results, and with less of the "personal element" than the volumetric.

Some modifications of the Fehling method, especially as carried out gravimetrically, differ for the various reducing sugars to be determined,

^{*} Leffmann and Beam, Select Methods of Food Analysis, p. 126.

and others are carried out alike, so far as manipulation is concerned, whether the particular sugar to be determined be dextrose, maltose, or lactose.

While, strictly speaking, the reducing power of dextrose, levulose, and invert sugar are not identical, it is customary in commercial work to regard them as such, and no appreciable error arises in consequence except in extreme cases. Thus the term "reducing sugars" is commonly applied indiscriminately to dextrose, levulose, and invert sugar, the same factor being used in calculating either, in mixtures wherein other reducing sugars, as lactose, maltose, etc., having widely different reducing powers are absent.

Fehling's solution is made up in two separate parts as follows:

- A. Fehling's Copper Solution.—34.639 grams of carefully selected crystals of pure copper sulphate dissolved in water and diluted to exactly 500 cc.
- B. Fehling's Alkaline Tartrate Solution.—173 grams Rochelle salts and 50 grams sodium hydroxide are dissolved in water and diluted to exactly 500 cc.

The Fehling solution should be standardized by dissolving 0.5 gram of pure anhydrous dextrose in water, and diluting to exactly 100 cc. 10 cc. of this dextrose solution should exactly reduce the copper in 10 cc. of the Fehling (5 cc. each of solutions A and B) when conducted according to the volumetric process described below.

VOLUMETRIC FEHLING PROCESS.—For determining dextrose, levulose, or invert sugar in a raw or brown sugar, make a solution of the sugar of such a strength that an accurately weighed amount dissolved in water and made up to 100 cc. shall contain not more than 1% of the reducing sugar, as nearly as can be guessed at with reference to the class of sugar under examination, or from a rough preliminary titration.

Measure accurately into a flask of about 250 cc. capacity 5 cc. Fehling's copper sulphate solution, A, and 5 cc. of the alkali solution, B. Add about 40 cc. of water, mix and boil over a free flame, with copper gauze beneath the flask. While still boiling, add from a pipette or burette a measured quantity of the sugar solution, prepared as above, until the copper after three minutes' boiling is all reduced to cuprous oxide. The end-point is determined in a variety of ways. Practice will soon enable the eye to judge the near approach of the end-point by the changes in color that take place in the solution, which turns from a deep blue, first to green, then to a dull-red tint, and finally to a bright brick-red. The sugar-containing solution may be added from the burette quite rapidly until

the solution reaches the dull-red tint, after which care is taken to add a little at a time, keeping account of the total amount added. If the flask be removed from the flame, and the bright, diffused light from a window viewed through the solution with the eye on a level with the surface, a thin film scarcely wider than a line will be observed just below the surface (see Fig. 109), which is blue so long as some of the copper in the solu-

tion remains unreduced. When, however, all the copper has been reduced, this film ceases to be blue and becomes colorless or yellow.

If the film is not at once apparent, it may often be made quite noticeable by simply diluting the solution in the flask with water. At the approach of the end-point the sugar-containing solution should be added a very little at a time. The exact end-point is best arrived at by decanting a few drops of the mixture in the flask through a filter, acidifying the filtrate with acetic acid, and adding a drop of a solution of ferrocyanide of potassium. As long as there is unreduced copper present, a precipitate or brown-red coloration will appear when the ferrocyanide is added. sugar solution toward the end should be added to the contents in the flask in small installments (say half a cubic centimeter each time), boiling the liquor for at least three minutes after each addition, until no brown-red coloration is pro-



Fig. 109.—Flask and Contents used in Volumetric Fehling Determinations. Showing layer just beneath the surface, the color of which indicates the end-point in adding the sugar-containing liquid.

duced by adding the ferrocyanide to a little of the filtered acidified liquid. When the number of cubic centimeters of sugar solution necessary to reduce the copper has thus been determined, a second titration should be made to verify the first, running the entire amount of sugar-containing liquid found necessary in the first case into the second flask.

The equivalents of 10 cc. of Fehling's solution in the above volumetric method are, in terms of the common reducing sugars, as follows:

o.0807 gram of maltose will reduce 10 cc. Fehling's solution.
o.067 gram of lactose "" 10 cc. ""

Suppose, for example, a sample of brown sugar is to be examined for invert sugar. This class of sugar has usually from 2 to 6 per cent of invert sugar. Hence, if 10 grams of the sample are dissolved in 100 cc., the resulting solution will contain not more than 1% of invert sugar.

Suppose 12.9 cc. of this 10% sugar solution were found by the above process to reduce 10 cc. of Fehling's solution.

10 cc. Fehling's solution are equivalent to 0.05 gram invert sugar.

Therefore 12.9 cc. of the sugar solution contain 0.05 gram invertsugar.

100 cc. sugar solution contain 10 grams sample, and 12.9 cc. contain 1.29 grams sample, the equivalent of 0.05 gram invert sugar.

Hence per cent invert sugar =
$$\frac{0.05 \times 100}{1.29}$$
 = 3.9.

GRAVIMETRIC FEHLING PROCESSES.—In determining reducing sugars by gravimetric processes, a measured volume of the sugar solution is allowed to act upon a measured volume of hot Fehling's solution for a fixed time, thus forming cuprous oxide. This may be dried and weighed direct, but is more commonly converted either into cupric oxide by ignition, or into metallic copper by reduction with hydrogen or by electrolysis. In any case the sugar is calculated from the weight of the cuprous oxide, the cupric oxide, or the metallic copper (whichever method be used) by the employment of the proper factor, or by the use of tables compiled for the purpose.

Note.—Much difference of opinion exists as to the best and most accurate Fehling gravimetric method to employ. For the determination of dextrose, the Association of Official Agricultural Chemists has given its approval to the Allihn method, wherein the cuprous oxide deposited is further reduced to metallic copper and the dextrose calculated from the copper by Allihn's table.

The author for two reasons prefers the method of O'Sullivan as employed by Defren, with the use of the Defren tables, in accordance with which the reducing sugar is expressed in terms of its equivalence to cupric oxide, first because of its comparative simplicity, involving as it does less processes than the Allihn method (each additional process introducing a possible source of error), and, second, because the same method as carried out is applicable for the determination not only of dextrose, but also of maltose and lactose, Defren having worked out

tables adapted for them all. Munsen and Walker* have also devised a simple method with accompanying tables, adapted, with a uniform system of procedure, to the determination of the various reducing sugars. In using the tables for dextrose, maltose, and lactose compiled by Allihn, Wein, and Soxhlet, the method employed must in each case be carried out in strict accordance to the minutest details adopted by each of the above authorities, and they are by no means uniform.

The Defren-O'Sullivan Method.†—Mix 15 cc. of Fehling's copper solution, A (p. 591), with 15 cc. of the tartrate solution, B, in a quarter-liter Erlenmeyer flask, and add 50 cc. of distilled water. Place the flask and its contents in a boiling water bath and allow them to remain five minutes. Then run rapidly from a burette into the hot liquor in the flask 25 cc. of the sugar solution to be tested (which should contain not more than one-half per cent of reducing sugar). Allow the flask to remain in the boiling water bath just fifteen minutes after the addition of the sugar solution, remove, and with the aid of a vacuum filter the contents rapidly in a platinum or porcelain Gooch crucible containing a layer of prepared asbestos fiber about 1 cm. thick, the Gooch with the asbestos having been previously ignited, cooled, and weighed. The cuprous oxide precipitate is thoroughly washed with boiling distilled water till the water ceases to be alkaline.

The asbestos used should be of the long-fibered variety, and should be specially prepared as follows: Boil first with nitric acid (specific gravity 1.05 to 1.10), washing out the acid with hot water, then boil with a 25% solution of sodium hydroxide, and finally wash out the alkali with hot water. Keep the asbestos in a wide-mouthed flask or bottle, and transfer it to the Gooch by shaking it up in the water and pouring it quickly into the crucible while under suction.

Dry the Gooch with its contents in the oven, and finally heat to dull redness for fifteen minutes, during which the red cuprous oxide is converted into the black cupric oxide. If a platinum Gooch is used (and this variety is preferred by the writer), it may be heated directly over the low flame of a burner. If the Gooch is of porcelain, considerable care must be taken to avoid cracking the crucible, the heat being increased cautiously and the operation preferably conducted in a radiator or muffle. After oxidation as above, the crucible is transferred to a desiccator, cooled, and quickly weighed. From the milligrams of cupric oxide, calculate the milligrams of dextrose from the following table:

^{*} U. S. Dept. of Agric., Bur. of Chem., Bul. 107 (rev.), p. 241.

[†] Jour. Am. Chem. Soc., 18, 1896, p. 749, and Tech. Quart., 10, 1897, p. 167.

DEFREN'S TABLE FOR THE DETERMINATION OF DEXTROSE, MALTOSE, AND LACTOSE.

			און שוו	CTOSE.			
Milligrams of Cupric Oxide.	Milligrams of Dextrose.	Milligrams of Maltose.	Milligrams of Lactose.	Milligrams of Cupric Oxide.	Milligrams of Dextrose.	Milligrams of Maltose.	Milligrams of Lactose.
30	13.2	21.7	18.8	80	35-4	58.1	50.5
31	13.7	22.4	19.5	81	35-9	58.9	51.1
32	14.1	23.1	20.1	82	35.9	59.6	51.7
	14.6		20.7	83	36.3 36.8	59.0	
33		23.9		84		60.3 61.1	52.4
34	15.0	24.6	21.4	•	37 - 2	01.1	53.0
35	15.4	25.3	22.0	8 ₅ 86	37 - 7	61.8	53.6
36	15.9	26.1	22.6		38.1	62.5	54-3
37	16.3	26.8	23.3	87	38.5	63.3	54.9
38	16.8	27.5	23.9	88	39.0	64.0	55.5
39	17.2	28.3	24.5	89	39-4	64.7	56.2
40	17.6	29.0	25.2	90	39-9	65.5	56.8
41	18.1	29.7	25.8	91	40.3	66.2	57-4
.42	18.5	30.5	26.4	92	40.8	66.9	58.1
43	19.0	31.2	27.1	93	41.2	67.7	58.7
44	19.4	31.9	27.7	94	41.7	68.4	59-3
45	19.9	32.7	28.3	95	42.1	69.1	60.0
46	20.3	33-4	29.0	96	42.5	69.9	60.6
47	20.7	34.1	29.6	97	43.0	70.6	61.2
48	21.2	34.8	30.2	98	43-4	71.3	61.g
49	21.6	35.5	30.8	99	43.9	72.1	62.5
50	22.1	36.2	31.5	100	44-4	72.8	63.2
51	22.5	37.0	32.1	101	44.8	73.5	63.8
52	23.0	37.7	32.7	102	45.3	74-3	64.4
	23.4	38.4		103		75.0	65.1
53	23.8	39.2	33-3	103	45.7 46.2		65.7
54	-3.0	39.2	34.0	104	40.2	75-7	3.7
55	24.2	39-9	34.6	105	46.6	76.5	66.3
56	24-7	40-5	35.2	106	47.0	77.2	67.0
57	25.1	41.3	35-9	107	47-5	77-9	67.6
58	25-5	42.1	36.5	108	48.0	78.7	68.2
59	26.0	42.8	37.1	109	48.4	79-4	68.9
60	26.4	43.5	37.8	110	48.9	80.1	69.5
61	26.9	44-3	38.4	111	49.3	80.0	70.1
62	27.3	45.0	39.0	112	49.8	81.6	70.8
63	27.8	45.7	39.7	113	50.2	82.3	71.4
64	28.2	46.5	40.3	114	50.7	83.1	72.0
65	28.7	47-2	40.9	775		83.8	72.7
66	20.7	47.9	41.6	115	51.1	84.5	73-3
67		47.9		117		85.2	74.0
68	29.5		42.2	117	52.0	85.9	74.6
69	30.0 30.4	49·4 50.1	42.8 43-5	119	52.4 52.9	86.6	75.2
-9	3-14		43-3] 3		''
70	30.9	50.8	44-1	120	53-3	87.4 88.1	75-9 76.6
71	31.3	51.6	44-7	121	53.8	88.9	
72	31.8	52-3	45.4	122	54.2		77.2
73 74	32.2 32.6	53.0	46.0 46.6	123	54-7 55-1	89.6 90.3	77.9 78.5
•	3-1-	1	'				' -
75	33.1	54.5	47.3	125	55.6	91.1	79.1
76	33-5	55-2	47-9	126	56.0	91.8	79.8
77	34.0	56.0	48.5	127	56.5	92.5	80.4
78	34-4	56.7	49.2	128	56.9	93-3	81.1
79	34-9	57-4	49.8	11 129	57-3	94.0	81.7

DEFREN'S TABLE FOR THE DETERMINATION OF DEXTROSE, MALTOSE, AND LACTOSE—(Continued).

		AND	LACTOS	E—(Contin	rued).		
Milligrams of Cupric Oxide.	Milligrams of Dextrose.	Milligrams of Maltose.	Milligrams of Lactose.	Milligrams of Cupric Oxide.	Milligrams of Dextrose.	Milligrams of Maltose.	Milligrams of Lactose.
130	57.8	94.8	82.4	180	80.4	131.8	114.6
131	58.2	, ,	83.0	181	80.8		
132	58.7	95·5 96.2	83.6	182	8	132.5	115.2
		,	03.0		81.3 81.8	133.2	115.8
133	59.1	97.0	84.2	183		134.0	116.5
134	59.6	97 - 7	84.9	184	82.2	134-7	117.1
135	60.0	98.4	85.5	185	82.7	135.5	117.8
136	60.5	99.2	86.1	186	83.1	136.2	118.4
137	60.9	- 99.9	86.8	187	83.5	136.9	119.1
138	61.3	100.7	87.4	188	84.0	137.7	119.7
139	61.8	101.4	88.1	189	84.4	138.4	120.4
140	62.2	102.1	88.7	190	84.9	139.1	121.0
141	62.7	102.8	89.3	191	85.4	139.9	121.7
142	63.1	103.5	90.0	192	85.9	140.6	122.3
143	63.6	104.3	90.6	193	86.3	141.4	123.0
144	64.0	105.0	91.3	194	86.3 86.8	142.1	123.6
145	64.5	105.8	91.9	195	87.2	142.8	124.3
146	64.9	106.5	92.6	196	87.7	143.6	
147	65.4	107.2	93.2	197	88.1	143.0	124.9 125.6
148	65.8	108.0	93.9	198	88.6	145.1	125.0
149	66.3	108.7	94-5	199	89.0	145.8	126.9
150	66.8	****	0		0		
151	67.3	109.5	95.2	200 20I	89.5	146.6	127.5
152		111.0	95.8	201	89.9	147.3	128.2
	67.7 68.3		96.5	1	90.4	148.1	128.8
153	68.7	111.7	97.1	203	90.8	148.8	129.5
154	00.7	112.4	97.8	204	91.3	149.6	130.1
155	69.2	113.2	98.4	205	91.7	- 150.3	130.8
156	69.6	113.9	99.1	206	92.2	151.1	131.5
157	70.0	114-7	99 - 7	207	92.6	151.8	132.1
158	70-5	115.4	100.4	208	93.1	152.5	132.8
159	70.9	116.1	101.0	209	93-5	153-3	133-4
160	71.3	116.9	101.7	210	94.0	154.1	134.1
161	71.8	117.6	102.3	211	94-4	154.8	134.7
162	72.3	118.4	103.0	212	94-9	155.6	135.4
163	72.7	119.1	103.6	213	95-3	156.3	136.0
164	73-2	119.9	104.3	214	95.8	157.1	136.7
165	73.6	120.6	104.9	215	96.3	157.8	137-3
166	74-1	121.4	105.6	216	96.7	158.6	138.0
167	74-5	122.1	106.2	217	97.2	159.3	138.6
168	74-9	122.9	106.9	218	97.6	160.0	139.3
169	75-4	123.6	107.5	219	98.1	160.8	139.9
170	75.8	124.4	108.2	220	98.6	161.5	140.6
171	76.3	125.1	108.8	211	99.0	162.3	141.2
172	76.8	125.8	109.5	222	99.5	163.0	141.9
173	77-3	126.6	110.1	223	99.9	163.7	142.5
174	77.7	127.3	110.8	224	100.4	164.5	143.2
175	78.2	128.1	,,, ,	225	700.0		
175	78.6	128.8	111.4	225 226	100.9	165.3	143.8
177	79.1	1	112.6) I	101.3	166.0	144.5
178		129.5		227	101.8	166.8	145.1
179	79-5 80.0		113.3	1	102.2	167.5	145.8
1/9	, 60.0	131.0	113.9	229	102.7	168.3	146.4

DEFREN'S TABLE FOR THE DETERMINATION OF DEXTROSE, MALTOSE, AND LACTOSE—(Concluded).

			LACIOS	E-(Concil			,
Milligrams of Cupric Oxide,	Milligrams of Dextrose.	Milligrams of Maltose.	Milligrams of Lactose.	Milligrams of Cupric Oxide.	Milligrams of Dextrose.	Milligrams of Maltose.	Milligrams of Lactose,
230	103.1	169.1	147.0	280	126.1	206.8	179.6
231	103.6	169.8	147-7	281	126.5	207.5	180.2
		170.6	148.3	282	127.0	208.3	180.0
232	104.0			283			181.5
233	104.5	171.3	149.0		127.4	209.0	
234	105.0	172.1	149.6	284	127.9	209.8	182.2
235	105.4	172.8	150.3	285	128.3	210.5	182.9
236	105.9	173.6	150.9	286	128.8	211.3	183.6
237	106.3	174-3	151.6	287	129.3	212.1	184.2
238	106.8	175.1	152.2	288	129.7	212.8	184.9
239	107.2	175.8	152.9	289	130.2	213.6	185.6
240	107.7	176.6	153.5	290	130.6	214.3	186.2
241	108.1	177-3	154.2	291	131.1	215.1	186.g
242	108.6	178.1	154.8	202	131.5	215.9	187.6
243	100.0	178.8	135.5	293	132.0	216.6	188.2
	109.5	179.6	156.1	294	132.5	217.4	188.9
244	109.5	1/9.0	130.1	294	132.3	/	100.9
245	109.9	180.3	156.8	295	133.0	218.2	189.5
246	110.4	181.1	157.4	296	133-4	218.9	190.2
247	110.9	181.8	158.1	297	133.9	219.7	190.8
248	111.3	182.6	158.7	298	134.3	220.4	191.5
2 49	111.8	183.3	159-4	299	134.8	221.2	192.1
250	112.3	184.1	160.0	300	135-3	221.9	192.8
251	112.7	184.8	160.7	301	135.7	222.7	193.4
252	113.2	185.5	161.3	302	136.2	223.5	194.1
253	113.7	186.3	162.0	303	136.6	224.2	194.7
254	114.1	187.1	162.6	304	137.1	225.0	195.3
		187.8	762.2	205	7.77 6	225.8	196.0
255	114.6		163.3	305	137.6		
256	115.0	188.6	163.9	306	138.0	226.5	196.6
257	115.5	189.3	164.6	307	138.5	227.3	197-3
258	116.0	190.1	165.2	308	138.9	228.1	197.9
259	116.4	190.8	165.9	309	139.4	228.8	198.6
260	116.9	191.6	166.5	310	139.9	229.6	199.3
261	117.3	192.4	167.2	311	140.3	230.4	199.9
262	117.8	193.1	167.8	312	140.8	231.1	200.6
263	118.3	193.9	168.1	313	141.2	231.0	201.3
264	118.7	194.6	169.5	314	141.7	232.7	202.0
265	119.2	195.4	169.8	315	142.2	233.4	202.6
266	119.6	195.1	170.4	316	142.6	234-2	203.3
267	120.1	196.9	171.1	317	143.1	234.9	203.9
268	120.6		, ,	318	143.6	235.7	204.6
269	121.0	197.7	171.7	319	144.0	236.5	205.3
•	Ì	1	1	1		1	207.0
270	121.4	199.2	173.0	320	144-5	237.2	205.9
271	121.9	199.9	173-7	11		}	1
272	122.4	200.7	174-4	11	l		1
273 274	122.8	201.5	175.0		1		l
274	125.3	202.2	175-7	il	\	l	
275	123.7	203.0	176.3		I		1
276	124.2	203.7	177.0	!!	I	i	1
277	124.6	204.5	177.6		1	1	1
278	125.1	205.2	178.3	Į).	1	1	1
279	125.6	206.0	178.0	l,		1	l

Munson and Walker Method.*—1. Preparation of Solutions and Asbestos.—Use the copper sulphate solution and alkaline tartate solution as given on page 591. Prepare the asbestos, which should be the amphibole variety, by first digesting with 1:3 hydrochloric acid for two or three days. Wash free from acid, and digest for a similar period with soda solution, after which treat for a few hours with hot alkaline copper tartrate solution of the strength employed in sugar determinations. Then wash the asbestos free from alkali, finally digest with nitric acid for several hours, and after washing free from acid, shake with water for use. In preparing the Gooch crucible, load it with a film of asbestos one-fourth inch thick, wash this thoroughly with water to remove fine particles of asbestos; finally wash with alcohol and ether, dry for thirty minutes at 100° C., cool in a desiccator and weigh. It is best to dissolve the cuprous oxide with nitric acid each time after weighing, and use the same felts over and over again, as they improve with use.

2. Process.—Transfer 25 cc. each of the copper and alkaline tartrate solutions to a 400-cc. Jena or Non-sol beaker, and add 50 cc. of reducing sugar solution, or, if a smaller volume of sugar solution be used, add water to make the final volume 100 cc. Heat the beaker upon an asbestos gauze over a Bunsen burner, so regulate the flame that boiling begins in four minutes, and continue the boiling for exactly two minutes. Keep the beaker covered with a watch-glass throughout the entire time of heating. Without diluting, filter the cuprous oxide at once on an asbestos felt in a porcelain Gooch crucible, using suction. Wash the cuprous oxide thoroughly with water at a temperature of about 60° C., then with 10 cc. of alcohol, and finally with 10 cc. of ether. Dry for thirty minutes in a water oven at 100° C., cool in a desiccator and weigh as cuprous oxide.

The number of milligrams of copper reduced by a given amount of reducing sugar differs when sucrose is present and when it is absent. In the tables on pp. 599 to 607 the absence of sucrose is assumed, except in the two columns under invert sugar, where one for mixtures of invert sugar and sucrose (0.4 gram of total sugar in 50 cc. of solution), and one for invert sugar and sucrose when the 50 cc. of solution contains 2 grams of total sugar are given, in addition to the column for invert sugar alone.

^{*} Jour. Am. Chem. Soc., 28, 1906, p. 163; 29, 1907, p. 541; U. S. Dept. Agric., Bur. of Chem., Bul. 107 (rev.), p. 241.

					eights in	milligran	1S.]				
á				Invert and St	Sugar icrose.	1	Lactose.		Malt	ose.	ô
Cuprous Oxide (CusO).	Copper (Cu).	Dextrose.	Invert Sugar.	o.4 Gram Total Sugar.	2 Grams Total Sugar.	CuHwOn.	CuHmOu++HoO.	CızHmOıı+HrO.	CısH#On.	CuHaOu+HrO.	Cuprous Oxide (CusO)
10	8.9	4.0	4·5	1.6		3.8	3.9	4.0	5.9	6.2	10
11	9.8	4.5	5·0	2.1		4.5	4.6	4.7	6.7	7.0	11
12	10.7	4.9	5·4	2.5		5.1	5.3	5.4	7.5	7.9	12
13	11.5	5.3	5.8	3.0		5.8	5.9	6.1	8.3	8.7	13
14	12.4	5.7	6·3	3.4		6.4	6.6	6.8	9.1	9.5	14
15 16 17 18	13.3 14.2 15.1 16.0 16.9	6.2 6.6 7.0 7.5 7.9	6.7 7.2 7.6 8.1 8.5	3.9 4.3 4.8 5.2 5.7		7.1 7.7 8.4 9.1 9.7	7.3 8.0 8.6 9.3	7·5 8·2 8·8 9·5 10·2	9.9 10.6 11.4 12.2 13.0	10.4 11.2 12.0 12.9 13.7	15 16 17 18
20 21 22 23 24	17.8 18.7 19.5 20.4 21.3	8.3 8.7 9.2 9.6	8.9 9.4 9.8 10.3	6.1 6.6 7.0 7.5 7.9		10.4 11.0 11.7 12.3 13.0	10.6 11.3 12.0 12.7 13.3	10.9 11.6 12.3 13.0	13.8 14.6 15.4 16.2 17.0	14.6 15.4 16.2 17.1 17.9	20 21 22 23 24
25 26 27 28 29	22.2 23.1 24.0 24.9 25.8	10.5 10.9 11.3 11.8 12.2	11.2 11.6 12.0 12.5 12.9	8.4 8.8 9.3 9.7		13.6 14.3 15.0 15.6 16.3	14.0 14.7 15.3 16.0 16.7	14.4 15.1 15.7 16.4 17.1	17.8 18.6 19.4 20.2 21.0	18.7 19.6 20.4 21.2 22.1	25 26 27 28 29
30 31 32 33 34	26.6 27.5 28.4 29.3 30.2	12.6 13.1 13.5 13.9 14.3	13.4 13.8 14.3 14.7 15.2	10.7 11.1 11.6 12.0 12.5	4·3 4·7 5·2 5.6 6.1	16.9 17.6 18.2 18.9	17.4 18.0 18.7 19.4 20.1	17.8 18.5 19.2 19.9 20.6	21.8 22.6 23.3 24.1 24.9	22.9 23.7 24.6 25.4 26.2	30 31 32 33 34
35	31.1	14.8	15.6	12.9	6.5	20.2	20.7	21.3	25.7	27.1	35
36	32.0	15.2	16.1	13.4	7.0	20.9	21.4	22.0	26.5	27.9	36
37	32.9	15.6	16.5	13.8	7.4	21.5	22.1	22.7	27.3	28.7	37
38	33.8	16.1	16.9	14.3	7.9	22.2	22.8	23.3	28.1	29.6	38
39	34.6	16.5	17.4	14.7	8.4	22.8	23.4	24.0	28.9	30.4	39
40	35.5	16.9	17.8	15.2	8.8	23.5	24.1	24.7	29.7	31.3	40
41	36.4	17.4	18.3	15.6	9.3	24.1	24.8	25.4	30.5	32.1	41
42	37.3	17.8	18.7	16.1	9.7	24.8	25.4	26.1	31.3	32.9	42
43	38.2	18.2	19.2	16.6	10.2	25.4	26.1	26.8	32.1	33.8	43
44	39.1	18.7	19.6	17.0	10.7	26.1	26.8	27.5	32.9	34.6	44
45	40.0	19.1	20.1	17.5	11.1	26.8	27.5	28.2	33.7	35.4	45
46	40.9	19.6	20.5	17.9	11.6	27.4	28.1	28.8	34.4	36.3	46
47	41.7	20.0	21.0	18.4	12.0	28.1	28.8	29.5	35.2	37.1	47
48	42.6	20.4	21.4	18.8	12.5	28.7	29.5	30.2	36.0	37.9	48
49	43.5	20.9	21.9	19.3	12.9	29.4	30.1	30.9	36.8	38.8	49
50	44.4	21.3	22.3	19.7	13.4	30.0	30.8	31.6	37.6	39.6	50
51	45.3	21.7		20.2	13.9	30.7	31.5	32.3	38.4	40.4	51
52	46.2	22.2		20.7	14.3	31.3	32.1	33.0	39.2	41.3	52
53	47.1	22.6		21.1	14.8	32.0	32.8	33.6	40.0	42.1	53
54	48.0	23.0		21.6	15.2	32.6	33.5	34.3	40.8	42.9	54
55	48.9	23.5	24.6	22.0	15.7	33.3	34.1	35.0	41.6	43.8	55
56	49.7	23.9	25.0	22.5	16.2	33.9	34.8	35.7	42.4	44.6	56
57	50.6	24.3	25.5	22.9	16.6	34.6	35.5	36.4	43.2	45.4	57
58	51.5	24.8	25.9	23.4	17.1	35.2	36.1	37.1	44.0	46.3	58
59	52.4	25.2	26.4	23.9	17.5	35.9	36.8	37.7	44.8	47.1	59
60	53.3	25.6	26.8	24.3	18.0	36.5	37.5	38.4	45.6	48.0	60
61	54.2	26.1	27.3	24.8	18.5	37.2	38.2	39.1	46.3	48.8	61
62	55.1	26.5	27.7	25.2	18.9	37.8	38.8	39.8	47.1	49.6	62
63	56.0	27.0	28.2	25.7	19.4	38.5	39.5	40.5	47.9	50.5	63
64	56.8	27.4	28.6	26.2	19.8	39.2	40.2	41.2	48.7	51.3	64

MUNSON AND WALKER'S TABLE FOR CALCULATING DEXTROSE, INVERT SUGAR, LACTOSE, AND MALTOSE—(Continued).

[Weights in milligrams.]

-	· · · ·				eights in	mungra	ms. j				,
©				Invert and St			Lactose.		Mal	tose.	Q.
Cuprous Oxide (Cu2O).	Copper (Cu).	Dextrose.	Invert Sugar.	o.4 Gram Total Sugar.	2 Grams Total Sugar.	CızHzzOıı.	C13H#O11 + 1 H1O.	CuHzOu+HzO.	CısH u Oıı.	C12H#O11 + H5O.	Cuprous Oxide (CurO.)
65	57.7	27.8	29.1	26.6	20.3	39.8	40.9	41.9	49.5	52.1	65
66	58.6	28.3	29.5	27.1	20.8	40.5	41.6	42.6	50.3	53.0	66
67	59.5	28.7	30.0	27.5	21.2	41.1	42.2	43.3	51.1	53.8	67
68	60.4	29.2	30.4	28.0	21.7	41.8	42.9	44.0	51.9	54.6	68
69	61.3	29.6	30.9	28.5	22.2	42.5	43.0	44.7	52.7	55.5	69
. 70	62.2	30.0	31.3	28.9	22.6	43.1	44.3	45.4	53 · 5		70
71	63.1	30.5	31.8	29.4	23.1	43.8	44.9	46.1	54 · 3		71
72	64.0	30.9	32.3	29.8	23.5	44.4	45.6	46.8	55 · 1		72
73	64.8	31.4	32.7	30.3	24.0	45.1	40.3	47.5	55 · 9		73
74	65.7	31.8	33.2	30.8	24.5	45.7	47.0	48.2	56 · 7		74
75	66.6	32.2	33.6	31.2	24.9	46.4	47.6	48.8	57.5	60.5	75
76	67.5	32.7	34.1	31.7	25.4	47.0	48.3	49.5	58.2	61.3	76
77	68.4	33.1	34.5	32.1	25.9	47.7	49.0	50.2	59.0	62.1	77
78	69.3	33.6	35.0	32.6	26.3	48.4	49.6	50.9	59.8	63.0	78
79.	70.2	34.0	35.4	33.1	26.8	49.0	50.3	51.6	60.6	63.8	79
80	71.1	34.4	35.9	33.5	27.3	49.7	51.0	52.3	61.4	64.6	80
81	71.9	34.9	36.3	34.0	27.7	50.3	51.6	53.0	62.2	65.5	81
82	72.8	35.3	36.8	34.5	28.2	51.0	52.3	53.7	63.0	66.3	82
83	73.7	35.8	37.3	34.9	28.6	51.6	53.0	54.4	63.8	67.1	83
84	74.6	36.2	37.7	35.4	29.1	52.3	53.7	55.0	64.6	68.0	84
85	75.5	36.7	38.2	35.8	29.6	52.9	54.3	55.7	65.4	68.8	85
86	76.4	37.1	38.6	36.3	30.0	53.6	55.0	56.4	66.2	69.7	86
87	77.3	37.5	39.1	36.8	30.5	54.3	55.7	57.1	67.0	70.5	87
88	78.2	38.0	39.5	37.2	31.0	54.9	56.4	57.8	67.8	71.3	88
89	79.1	38.4	40.0	37.7	31.4	55.6	57.0	58.5	68.5	72.2	89
90	79.9	38.9	40.4	38.2	31.9	56.2	57.7	59.2	69.3	73.0	90
91	80.8	39.3	40.9	38.6	32.4	56.9	58.4	59.9	70.1	73.8	91
92	81.7	39.8	41.4	39.1	32.8	57.5	59.0	60.6	70.9	74.7	92
93	82.6	40.2	41.8	39.6	33.3	58.2	59.7	61.3	71.7	75.5	93
94	83.5	40.6	42.3	40.0	33.8	58.8	60.0	61.9	72.5	76.3	94
95	84.4	41.1	42.7	40.5	34.2	59.5	61.1	62.6	73.3	77.2	95
96	85.3	41.5	43.2	41.0	34.7	60.2	61.7	63.3	74.1	78.0	96
97	86.2	42.0	43.7	41.4	35.2	60.8	62.4	64.0	74.9	78.8	97
98	87.1	42.4	44.1	41.9	35.6	61.5	63.1	64.7	75.7	79.7	98
99	87.9	42.9	44.6	42.3	36.1	62.1	63.8	65.4	76.5	80.5	99
100	88.8	43.3	45.0	42.8	36.6	62.8	64.4	66.1	77.3	81.3	100
101	89.7	43.8	45.5	43.3	37.0	63.4	65.1	66.8	78.1	82.2	101
102	90.6	44.2	46.0	43.8	37.5	64.1	65.8	67.5	78.8	83.0	102
103	91.5	44.7	46.4	44.2	38.0	64.7	66.4	68.1	79.6	83.8	103
104	92.4	45.1	46.9	44.7	38.5	65.4	67.1	68.8	80.4	84.7	104
105	93.3	45.5	47.3	45.2	38.9	66.1	67.8	69.5	81.2	85.5	105
106	94.2	46.0	47.8	45.6	39.4	66.7	68.5	70.2	82.0	86.3	106
107	95.0	46.4	48.3	46.1	39.9	67.4	69.1	70.9	82.8	87.2	107
108	95.9	46.9	48.7	46.6	40.3	68.0	69.8	71.6	83.6	88.0	108
109	96.8	47.3	49.2	47.0	40.8	68.7	70.5	72.3	84.4	88.8	109
110	97.7	47.8	49.6	47.5	41.3	69.3	71.1	73.0	85.2	89.7	110
111	98.6	48.2	50.1	48.0	41.7	70.0	71.8	73.6	86.0	90.5	111
112	99.5	48.7	50.6	48.4	42.2	70.6	72.5	74.3	86.8	91.3	112
113	100.4	49.1	51.0	48.9	42.7	71.3	73.1	75.0	87.6	92.2	113
114	101.3	49.6	51.5	49.4	43.2	71.9	73.8	75.7	88.4	93.0	114
115	102.2	50.0	51.9	49.8	43.6	72.6	74-5	76.4	89.2	93·9	115
116	103.0	50.5	52.4	50.3	44.1	73.2	75-2	77.1	90.0	94·7	116
117	103.9	50.9	52.9	50.8	44.6	73.9	75.8	77.8	90.7	95·5	117
118	104.8	51.4	53.3	51.2	45.0	74.5	76.5	78.5	91.5	96·4	118
119	105.7	51.8	53.8	51.7	45.5	75.2	77.2	79.1	92.3	97·2	119

				Į W	eights in	mungran	09.]				
ĝ				Invert and St	Sugar icrose.		Lactose.		Malt	ose.	ું
Cuprous Oxide (CusO).	Copper (Cu).	Dextrose.	Invert Sugar.	o.4 Gram Total Sugar.	2 Grams Total Sugar.	CısH#Ou.	CuH2Ou+\$H2O.	CaH#O11 + HsO	СыНжОп.	CuHzOn + H4O.	Cuprous Oxide (CusO).
120 121 122 123 124	106.6 107.5 108.4 109.3	52.3 52.7 53.2 53.6 54.1	54·3 54·7 55·2 55·7 56·1	52.2 52.7 53.1 53.6 54.1	46.0 46.5 46.9 47.4 47.9	75.8 76.5 77.1 77.8 78.5	77.8 78.5 79.2 79.9 80.5	79.8 80.5 81.2 81.9 82.6	93.1 93.9 94.7 95.5 96.3	98.0 98.9 99.7 100.5	120 121 122 123 124
125 126 127 128 129	111.0 111.9 112.8 113.7 114.6	54.5 55.0 55.4 55.9 56.3	56.6 57.5 57.5 58.0 58.4	54.5 55.0 55.5 55.9 56.4	48.3 48.8 49.3 49.8 50.2	79.1 79.8 80.4 81.1 81.7	81.2 81.9 82.5 83.2 83.9	83 · 3 84 · 0 84 · 7 85 · 4 86 · 0	97.1 97.9 98.7 99.4 100.2	102.2 103.0 103.9 104.7 105.5	125 126 127 128 129
130 131 132 133 134	115.5 116.4 117.3 118.1 119.0	56.8 57.2 57.7 58.1 58.6	58.9 59.4 59.8 60.3 60.8	56.9 57.4 57.8 58.3 58.8	50.7 51.2 51.7 52.1 52.6	82.4 83.1 83.7 84.4 85.0	84.6 85.2 85.9 86.6 87.3	86.7 87.4 88.1 88.8 89.5	101.0 101.8 102.6 103.4 104.2	106.4 107.2 108.0 108.9 109.7	130 131 132 133 134
135 136 137 138 139	119.9 120.8 121.7 122.6 123.5	59.0 59.5 60.0 60.4 60.9	61.2 61.7 62.2 62.6 63.1	59.3 59.7 60.2 60.7 61.2	53.1 53.6 54.0 54.5 55.0	85.7 86.3 87.0 87.7 88.3	87.9 88.6 89.3 90.0 90.6	90.2 90.9 91.6 92.3 93.0	105.0 105.8 106.6 107.4 108.2	110.5 111.4 112.2 113.0 113.9	135 136 137 138 139
140 141 142 143 144	124.4 125.2 126.1 127.0 127.9	61.3 61.8 62.2 62.7 63.1	63.6 64.0 64.5 65.0 65.4	61.6 62.1 62.6 63.1 63.5	55.5 55.9 56.4 56.9 57.4	89.0 89.6 90.3 90.9 91.6	91.3 92.0 92.6 93.3 94.0	93.6 94.3 95.0 95.7 96.4	109.0 109.8 110.5 111.3	114.7 115.5 116.4 117.2 118.0	140 141 142 143 144
145 146 147 148 149	128.8 129.7 130.6 131.5 132.4	63.6 64.0 64.5 65.0 65.4	65.9 66.4 66.9 67.3 67.8	64.0 64.5 65.0 65.4 65.9	57.8 58.3 58.8 59.3 59.7	92.2 92.9 93.5 94.2 94.8	94.7 95.3 96.0 96.7 97.3	97.1 97.8 98.4 99.1 99.8	112.9 113.7 114.5 115.3 116.1	118.9 119.7 120.5 121.4 122.2	145 146 147 148 149
150 151 152 153 154	133.2 134.1 135.0 135.9 136.8	65.9 66.3 66.8 67.2 67.7	68.3 68.7 69.2 69.7 70.1	66.4 66.9 67.3 67.8 68.3	60.2 60.7 61.2 61.7 62.1	95.5 96.2 96.8 97.5 98.1	98.0 98.7 99.3 100.0	100.5 101.2 101.9 102.6 103.3	116.9 117.7 118.5 119.3 120.0	123.0 123.9 124.7 125.5 126.4	150 151 152 153 154
155 156 157 158 159	137.7 138.6 139.5 140.3 141.2	68.2 68.6 69.1 69.5 70.0	70.6 71.1 71.6 72.0 72.5	68.8 69.2 69.7 70.2 70.7	62.6 63.1 63.6 64.1 64.5	98.8 99.4 100.1 100.7 101.4	101.4 102.0 102.7 103.4 104.1	104.0 104.7 105.3 106.0 106.7	120.8 121.6 122.4 123.2 124.0	127.2 128.0 128.9 129.7 130.5	155 156 157 158 159
160 161 162 163 164	142.1 143.0 143.9 144.8 145.7	70.4 70.9 71.4 71.8 72.3	73.0 73.4 73.9 74.4 74.9	71.2 71.6 72.1 72.6 73.1	65.0 65.5 66.0 66.5 66.9	102.0 102.7 103.4 104.0	104.7 105.4 106.1 106.7 107.4	107.4 108.1 108.8 109.5	124.8 125.6 126.4 127.2 128.0	131.4 132.2 133.0 133.9 134.7	160 161 162 163 164
165 166 167 168 169	146.6 147.5 148.3 149.2 150.1	72.8 73.2 73.7 74.1 74.6	75.3 75.8 76.3 76.8 77.2	73.6 74.0 74.5 75.0 75.5	67.4 67.9 68.4 68.9 69.3	105.3 106.0 106.6 107.3 107.9	108.1 108.8 109.4 110.1 110.8	110.9 111.5 112.2 112.9 113.6	128.8 129.6 130.3 131.1 131.9	135.5 136.4 137.2 130.0 138.9	165 166 167 168 169
170 171 172 173 174	151.0 151.9 152.8 153.7 154.6	75.1 75.5 76.0 76.4 76.9	77.7 78.2 78.7 79.1 79.6	76.0 76.4 76.9 77.4 77.9	69.8 70.3 70.8 71.3 71.7	108.6 109.2 109.9 110.5	111.4 112.1 112.8 113.5 114.1	114.3 115.0 115.7 116.4 117.1	132.7 133.5 134.3 135.1 135.9	139.7 140.5 141.4 142.2 143.0	170 171 172 173 174

				(We	ights in	milligran	ns.]				
ĝ				Invert and Su			Lactose.		Malt	000.	ç
Cuprous Oxide (CusO).	Copper (Cu),	Dextrose.	Invert Sugar.	o.4 Gram Total Sugar.	2 Grams Total Sugar.	CatHatOu.	C14H23O11+ \$H5O.	C13H22O11 + H5O.	CuHnOu.	CuHaOu + HaO.	Cuprous Oxide (CusO)
175 176 177 178 179	155.5 156.3 157.2 158.1 159.0	77.4 77.8 78.3 78.8 79.2	80.1 80.6 81.0 81.5 82.0	78.4 78.8 79.3 79.8 80.3	72.2 72.7 73.2 73.7 74.2	111.9 112.5 113.2 113.8 114.5	115.5 1 116.1 1 116.8 1	17.7 18.4 19.1 19.8 20.5	136.7 137.5 138.3 139.1 139.8	143.9 144.7 145.5 146.4 147.2	175 176 177 178 179
180 181 182 183 184	159.9 160.8 161.7 162.6 163.4	79.7 80.1 80.6 81.1 81.5	82.5 82.9 83.4 83.9 84.4	80.8 81.3 81.7 82.2 82.7	74.6 75.1 75.6 76.1 76.6	115.1 115.8 116.5 117.1 117.8	118.8 I 119.5 I 120.2 I	21.2 21.9 22.6 23.3 23.9	140.6 141.4 142.2 143.0 143.8	148.0 148.9 149.7 150.5 151.4	180 181 182 183 184
185 186 187 188 189	164.3 165.2 166.1 167.0 167.9	82.0 82.5 82.9 83.4 83.9	84.9 85.3 85.8 86.3 86.8	83.2 83.7 84.2 84.6 85.1	77.1 77.6 78.0 78.5 79.0	118.4 119.1 119.7 120.4 121.0	123.5 1	24.6 25.3 26.0 26.7 27.4	144.6 145.4 146.2 147.0 147.8	152.2 153.0 153.9 154.7 155.5	185 186 187 188 189
190 191 192 193 194	168.8 169.7 170.5 171.4 172.3	84.3 84.8 85.3 85.7 86.2	87.2 87.7 88.2 88.7 89.2	85.6 86.1 86.6 87.1 87.6	79.5 80.0 80.5 81.0 81.4	121.7 122.3 123.0 123.6 124.3	125.5 1 126.2 1 126.9 1	28.1 28.8 29.5 30.1	148.6 149.3 150.1 150.9 151.7	156.4 157.2 158.0 158.9 159.7	190 191 192 193 194
195 196 197 198 199	173.2 174.1 175.0 175.9 176.8	86.7 87.1 87.6 88.1 88.5	90.6 90.6 91.1 91.6	88.0 88.5 89.0 89.5 90.0	81.9 82.4 82.9 83.4 83.9	125.6 125.6 126.3 126.9 127.6	128.9 1 129.6 1 130.3 1	31.5 32.2 32.9 33.6 34.3	152.5 153.3 154.1 154.9 155.7	160.5 161.4 162.2 163.0 163.9	195 196 197 198 199
200 201 202 203 204	177.7 178.5 179.4 180.3 181.2	89.0 89.5 89.9 90.4 90.9	92.0 92.5 93.0 93.5 94.0	90.5 91.0 91.4 91.9 92.4	84.4 84.8 85.3 85.8 86.3	128.2 128.9 129.5 130.2 130.8	132.3 132.9 133.6 134.3	35.0 35.7 36.3 37.0	156.5 157.3 158.1 158.8 159.6	164.7 165.5 166.4 167.2 168.0	200 201 202 203 204
205 206 207 208 209	182.1 183.0 183.9 184.8 185.6	91.4 91.8 92.3 92.8 93.2	94.5 94.9 95.4 95.9 96.4	92.9 93.4 93.9 94.4 94.9	86.8 87.3 87.8 88.3 88.8	131.5 132.1 132.8 133.4 134.1	135.6 136.3 137.0 137.6	138.4 139.1 139.8 140.5	160.4 161.2 162.0 162.8 163.6	168.9 169.7 170.5 171.4 172.2	205 206 207 208 209
210 211 212 213 214	186.5 187.4 188.3 189.2 190.1	93.7 94.2 94.6 95.1 95.6	96.9 97.4 97.8 98.3 98.8	95.4 95.8 96.3 96.8 97.3	89.2 89.7 90.2 90.7 91.2	134.8 135.4 136.1 136.7 137.4	139.6 139.6 140.3 141.0	141.9 142.5 143.2 143.9 144.6	164.4 165.2 166.0 166.8 167.5	173.0 173.8 174.7 175.5 176.4	210 211 212 213 214
215 216 217 218 219	191.0 191.9 192.8 193.6 194.5	96.1 96.5 97.0 97.5 98.0	99.3 99.8 100.3 100.8	97.8 98.3 98.8 99.3 99.8	91.7 92.2 92.7 93.2 93.7	138.0 138.7 139.3 140.0 140.6	142.3 143.0 143.7 144.3	145.3 146.0 146.7 147.3 148.0	168.3 169.1 169.9 170.7 171.5	177.2 178.0 178.9 179.7 180.5	215 216 217 218 219
220 221 222 223 224	195.4 :96.3 197.2 198.1 199.0	98.4 98.9 99.4 99.9 100.3	101.7 102.2 102.7 103.2 103.7	100.3 100.8 101.2 101.7 102.2	94.2 94.7 95.1 95.6 96.1	141.3 141.9 142.6 143.2 143.9	145.7 1 146.3 1 147.0 1 147.7 1	48.7 49.4 50.1 50.8 51.5	172.3 173.1 173.9 174.7 175.5	181.4 182.2 183.0 183.9 184.7	220 221 222 223 224
225 226 227 228 229	199 9 200.7 201.6 202.5 203.4	100.8 101.3 101.8 102.2 102.7	104.2 104.6 105.1 105.6 106.1	102.7 103.2 103.7 104.2 104.7	96.6 97.1 97.6 98.1 98.6	144.6 145.2 145.9 146.5 147.2	149.0 I 149.7 I 150.4 I	52.2 52.9 53.6 54.2 54.9	176.2 177.0 177.8 178.6 179.4	185.5 186.4 187.2 188.0 188.8	225 226 227 228 229

6				Invert and St	Sugar		Lactose.		Mai	tose.	Ĝ.
Cuprous Oxide (CusO)	Copper (Cu).	Dextrose.	Invert Sugar.	o.4 Gram Total Sugar.	s Grams Total Sugar.	CuH#On.	C13H22O11 + 1/2 H4O.	CuHmOn+HsO.	CuH#On.	CuHaOu+HaO.	Cuprous Oxide (CusO)
230	204.3	103.2	106.6	105.2	99.1	147.8	151.7	155.6	180.2	189.7	230
231	205.2	103.7	107.1	105.7	99.6	148.5	152.4	156.3	181.0	190.5	231
232	206.1	104.1	107.6	106.2	100.1	149.1	153.1	157.0	181.8	191.3	232
233	207.0	104.6	108.1	106.7	100.6	149.8	153.7	157.7	182.6	192.2	233
234	207.9	105.1	108.6	107.2	101.1	150.5	154.4	158.4	183.4	193.0	234
235 236 237 238 239	208.7 209.6 210.5 211.4 212.3	105.6 106.0 106.5 107.0	109.1 109.5 110.0 110.5	107.7 108.2 108.7 109.2 109.6	101.6 102.1 102.6 103.1 103.5	151.1 151.8 152.4 153.1 153.7	155.1 155.8 156.4 157.1 157.8	159.1 159.7 160.4 161.1 161.8	184.2 184.9 185.7 186.5 187.3	193.8 194.7 195.5 196.3 197.2	235 236 237 238 239
240	213.2	108.0	111.5	110.1	104.0	154.4	158.4	162.5	188.1	198.0	240
241	214.1	108.4	112.0	110.6	104.5	155.0	159.1	163.2	188.9	198.8	241
242	215.0	108.9	112.5	111.1	105.0	155.7	159.8	163.9	189.7	199.7	242
243	215.8	109.4	113.0	111.6	105.5	156.3	160.5	164.6	190.5	200.5	243
244	216.7	109.9	113.5	112.1	106.0	157.0	161.1	165.3	191.3	201.3	244
245 246 247 248 249	217.6 218.5 219.4 220.3 221.2	110.4 110.8 111.3 111.8 112.3	114.0 114.5 115.0 115.4 115.9	112.6 113.1 113.6 114.1 114.6	106.5 107.0 107.5 108.0 108.5	157.7 158.3 159.0 159.6 160.3	161.8 162.5 163.1 163.8 164.5	166.6 167.3 168.0 168.7	192.1 192.9 193.6 194.4 195.2	202.2 203.0 203.8 204.7 205.5	245 246 247 248 249
250 251 252 253 254	222.1 223.0 223.8 224.7 225.6	112.8 113.2 113.7 114.2 114.7	116.4 116.9 117.4 117.9 118.4	115.1 115.6 116.1 116.6 117.1	109.0 109.5 110.0 110.5	160.9 161.6 162.2 162.9 163.5	165.2 165.8 166.5 167.2 167.9	169.4 170.1 170.8 171.5 172.2	196.0 196.8 197.6 198.4 199.2	206.3 207.2 208.0 208.8 209.7	250 251 252 253 254
255	226.5	115.2	118.9	117.6	111.5	164.2	168.5	172.8	200.0	210.5	255
256	227.4	115.7	119.4	118.1	112.0	164.8	169.2	173.5	200.8	211.3	256
257	228.3	116.1	119.9	118.6	112.5	165.5	169.9	174.2	201.6	212.2	257
258	229.2	116.6	120.4	119.1	113.0	166.2	170.5	174.9	202.3	213.0	258
259	230.1	117.1	120.9	119.6	113.5	166.8	171.2	175.6	203.1	213.8	259
260 261 262 263 264	231.0 231.8 232.7 233.6 234.5	117.6 118.1 118.6 119.0	121.4 121.9 122.4 122.9 123.4	120.1 120.6 121.1 121.6 122.1	114.0 114.5 115.0 115.5 116.0	167.5 168.1 168.8 169.4 170.1	171.9 172.5 173.2 173.9 174.6	176.3 177.0 177.7 178.3 179.0	203.9 204.7 205.5 206.3 207.1	214.7 215.5 216.3 217.2 218.0	261 261 262 263 264
265	235.4	120.0	123.9	122.6	116.5	170.7	175.2	179.7	207.9	218.8	265
266	236.3	120.5	124.4	123.1	117.0	171.4	175.9	180.4	208.7	219.7	266
267	237.2	121.0	124.9	123.6	117.5	172.0	176.6	181.1	209.5	220.5	267
268	238.1	121.5	125.4	124.1	118.0	172.7	177.2	181.8	210.3	221.3	268
269	238.9	122.0	125.9	124.6	118.5	173.3	177.9	182.5	211.0	222.1	269
270	239.8	122.5	126.4	125.1	119.0	174.0	178.6	183.2	211.8	223.0	270
271	240.7	122.9	126.9	125.6	119.5	174.6	179.2	183.8	212.6	223.8	271
272	241.6	123.4	127.4	126.2	120.0	175.3	179.9	184.5	213.4	224.6	272
273	242.5	123.9	127.9	126.7	120.6	176.0	180.6	185.2	214.2	225.5	273
274	243.4	124.4	128.4	127.2	121.1	176.6	181.3	185.9	215.0	226.3	274
275	244.3	124.9	128.9	127.7	121.6	177.3	181.9	186.6	215.8	227.1	275
276	245.2	125.4	129.4	128.2	122.1	177.9	182.6	187.3	216.6	228.0	276
277	246.1	125.9	129.9	128.7	122.6	178.6	183.3	188.0	217.4	228.8	277
278	246.9	126.4	130.4	129.2	123.1	179.2	184.0	188.7	218.2	229.6	278
279	247.8	126.9	130.9	129.7	123.6	179.9	184.6	189.4	218.9	230.5	279
280	248.7	127.3	131.4	130.2	124.1	180.6	185.3	190.1	219.7	231.3	280
281	249.6	127.8	131.9	130.7	124.6	181.2	186.0	190.7	220.5	232.1	281
282	250.5	128.3	132.4	131.2	125.1	181.9	186.6	191.4	221.3	233.0	282
283	251.4	128.8	132.9	131.7	125.6	182.5	187.3	192.1	222.1	233.8	283
284	252.3	129.3	133.4	132.2	126.1	183.2	188.0	192.8	222.9	234.6	284

				[Weigi	its in mi	lligrama.	<u> </u>				
6				Invert and St	Sugar scrose.	<u> </u>	Lectose.		Mal	iose.	Ć
Cuprous Oxide (CusO).	Copper (Cu).	Dextruse.	Invert Sugar.	o.4 Gram Total Sugar	2 Grams Total Sugar.		CartinOta+\$HsO.	CısHaOıı + HaO.	CeHaOii.	CuHaOu + H4O.	Guprous Oxide (CurO).
285 286 287 288 289	253 2 254.0 254.9 255.8 250.7	129.8 130.3 130.8 131.3 131.8	133.9 134.4 134.9 135.4 135.9	132 7 133 2 133 7 134 3 134 8	126 6 127 1 127.6 128 1 128.6	183.8 184.5 185.1 185.8 186.4	188.7 189.3 190.0 190.7	193 5 194 2 194 9 195 5 196 2	223 7 224 5 225 3 226 1 226 9	235,5 236 3 237 1 238 0 238 8	285 286 287 288 288
290 291 292 293 294	257 6 258 5 259-4 260-3 261 2	132 3 132 7 133 2 133 7 134.2	136,4 136,9 137 4 137 9 138.4	135 3 135 8 136 3 136 8	129 2 129 7 130 2 130 7 131 2	187.1 187.7 188.4 189.0 189.7	192 0 192 7 193 3 194 0	196.9 197 6 198 3 199 0 199.7	227 6 228 4 229 2 230 0 230 8	239.6 240.5 241.3 242 1 242 9	290 291 292 293 294
295 296 297 298 299	262 0 262.9 263.8 264.7 265.6	134 7 135.2 135.7 136.2 136.7	138 9 139 4 140 0 140 5 141.0	137 8 138 3 138 8 139 4 139.9	131.7 132.2 132.7 133.7	190 3 191 0 191 7 192 3 193 0	195 4 196 0 196 7 197 4 198 0	200 .4 201 0 201 7 202 4 203 1	231 6 232 4 233 2 234 0 234 8	243 8 244 6 245 4 246 3 247 I	296 296 297 298 299
300 301 302 303 304	266 5 267 4 268 3 269 1 270 0	137 2 137.7 138 2 138 7 139 2	141.5 142.0 142.5 143.0 143.5	140 4 140 9 141 4 141 9 142 4	134.2 134.8 135.3 135.8 136.3	193 6 194 3 194 9 195 6 196 2	198 7 199 4 200 0 200 7 201 4	203 8 204 5 205 2 205 9 206 5	235.5 236 3 237 1 237 9 238 7	247 0 248 8 249 6 250.4 251.3	300 301 302 303 304
305 306 307 308 309	270 9 271 8 272 7 273.0 274 5	139 7 140 2 140 7 141 2 141 7	144 0 144 5 145 0 145 5 146 I	142 9 143 4 144 0 144 5 145 0	136 8 137 3 137 8 138 3 138 8	196 p 197 5 198 1 198 8 199 5	1 3 -	207 2 207 9 208 6 209 3 210.0	239 5 240 3 241 E 241 9 242 7	252.1 252.9 253.8 254.6 255.4	305 306 307 308 309
310 311 319 313 314	275.4 276 3 277.1 278 0 278 9	142 2 142 7 143 2 143 7 144 2	148 1	145.5 146 0 146 5 147 0	139.4 139.9 140.4 140.9 141.4	200 t 200 B 201 4 202 t 203 B	205 4 206 1 206 7 207 4 208.1	710.7 211 4 712.1 212 7 213.4	243 5 244 3 245 0 245 8 246 6	256 3 257 1 257 9 258 8 259 6	310 311 313 313 314
315 316 317 318 319	279 8 280 7 281 6 282.5 283 4	144 7 145.2 145 7 146 8 146.7	149 f 149 6 150 l 150.7 151 2	148 1 148 6 149 1 149 6 150 t	141 9 142 4 143 0 143 5 144 9	203.4 204 1 204 7 205 4 206 0	208 8 209 4 210 1 210 8 211 5	214.1 214 8 215 5 216 2 216 9	247 4 248 2 249 0 249 8 250 6	260 4 261 2 262.1 262.9 263.7	315 316 317 318 319
320 321 322 323 324	284 2 285.1 286 0 286.9 287 8	147 2 147.7 148 2 148.7 149 2	151 7 152.2 152 7 153.2 153 7	150 7 151.2 151.7 152.2 152.7	144 S 145 G 145 S 146 G 146 G	206 ? 207.3 208 0 208 6 209 3	212 I 212 8 213 5 214 I 214 8	217.6 218 3 218 9 219 6 220.3	251 3 252.1 252 9 253.7 254.5	264 6 265.4 266 2 267.1 267 9	320 321 328 323 324
325 320 327 328 329	288 7 289 6 290 5 291.4 292.2	149 7 150 8 150 7 151.2 351.7	154-3 154-8 155-3 155-8 150-3	153 2 153 8 154 3 154 8	147 t 147.6 148 t 148 6 149 t	210.6 210.6 211.3 211.9 312.6	215 5 216 2 216 8 217.5 218 2	221.0 221.7 222.4 223.1 223.8	255 3 256 1 256 9 257 7 258 5	2687 269 6 270.4 271 2 272.1	325 326 327 328 329
330 331 332 333 334	293 1 294 0 294 9 295.8 296 7	152.7 152.7 153.2 153.7 154.2	156 8 157 3 157.9 158.4 158.9	155 8 156 4 156 9 157 4 157 9	149.7 150 2 150 7 151 2 151 7	213 8 213 9 214 5 215 2 215 8	219 5 220 2 220 8	224 4 225 1 225 8 226 5 227 2	259 3 260 0 260.8 261 6 262 4	272 9 273 7 274.6 275 4 276 3	33° 33° 33° 333 334
335 336 337 338 339	297 6 298 5 299.3 300.2 301.1	154 7 155 2 155 8 150.3 156.8	159 4 159,9 160 5 161.0	158 4 159 0 159 5 169.0 160.5	152 B 152 B 153 3 153 B 154 3	216.5 217 1 217 8 218 4 219.1	222.9 222.9 223.5 224.2 224.9	227 9 228 6 229 2 229 9 230 6	264.0 264.0 264.8 265.6 266.4	277.0 277.9 278.7 279.5 280.4	335 336 337 338 339

MUNSON AND WALKER'S TABLE FOR CALCULATING DEXTROSE, INVERT SUGAR, LACTOSE, AND MALTOSE—(Continued).

[Weights in milligrams.]

						uningi si	ua. j				
Ĝ.					Sugar icrose.	:	Lactose,		Malt	ose.	ĝ
Cuprous Oxide (CurO).				Sugar.	s Grams Total Sugar.	CaHaOn.	Cathadu + \$Hio.	CtsHarOu + HaO.	CaHaOu.	CaHaOu+HaO.	Cuprous Oxide (CurO).
341 342 343 344	303.0 302.9 303 B 304 7 305.6	157-3 157-8 158-3 158-8 159-3	162.0 162.5 163 t 163 6 164 1	161 0 161 6 162 1 162 6 163 1	154 8 155 4 155 9 156 4 156 9	219.8 220.4 221.1 221.7 222.4	295.5 286 8 226 9 227 5 228 2	231.3 232.0 232.7 233.4 234.1	267.1 267.9 268.7 269.5 270.3	281.2 282.0 282.9 283.7 284.5	340 341 342 343 344
345 346 347 348 349	306.5 307.3 308.2 309.1 310.0	159 8 160.3 160.8 161 4 161 9	164 6 165 1 165.7 166.2 166 7	163.7 164.2 164.7 165.2 165.7	157 5 158.0 158 5 159.0 159 5	223 0 223 7 224.3 225.0 225.6	228.9 229 6 230 2 230.9 231.6	234.7 235.4 236.1 236.8 237.5	271.1 271.9 272.7 273.5 274.3	285 4 286 2 287 9 287 9 288 7	345 346 347 348 349
350 351 352 353 354	310.9 311 8 312.7 313 6 314.4	162 4 162 9 163 4 163-9 164 4	167 7 167.7 168 3 168.8 169 3	166 3 166 8 167 3 167.8 168 4	160 1 160.6 161 1 161.6 162 2	226 3 226.9 227.6 228 2 228.9	233.2 232 9 233 6 234 2 234 9	238.2 238.9 239.6 240.2 240.9	275 0 275 8 276 6 277.4 278 2	289.5 290.4 291.2 292.0 292.8	350 351 353 353 354
355 356 357 358 359	315,3 316,2 317,1 318 o 318 9	164 9 165 4 166 0 166 5	169 B 170.4 170 9 171.4 171 9	168 9 169.4 170.0 170.5 171.0	163.7 163.2 163.7 164.3 164.8	229.5 230 2 230.8 231 5 232 1	235.6 236.3 236.9 237.6 238.3	241.6 242 3 243 0 243.7 244.4	279 0 279 8 280 6 281 4 282 2	293 7 294.5 295.3 296.7 297 0	355 356 357 358 359
360 361 362 363 364	319.8 329 7 321.6 323 4 323.3	167.5 168.0 168.5 169.0 169.6	172.5 173.0 173.5 174.0 174.6	171.5 172.1 172.6 173.1 173.7	165.3 165.8 166.4 166.9	252.8 233 5 234 1 234 8 235.4	238.9 239 6 240.3 241 0 241.6	245.1 245.8 246.4 247.1 247.8	283 9 283 7 284 5 285 3 286 1	297.8 298 7 299.5 300.3 301.2	360 361 363 363 364
365 366 367 368 369	324.2 325.1 326.0 126.9 327.8	170 1 170 6 171 1 171.6 172.1	175 1 175 6 176 1 176 7 177 2	174-2 174-7 175-2 175-8 176-3	167 9 168 5 169 0 169.5 170 0	236 1 236 7 237.4 238 1 238.7	242 3 243.0 243 6 244 3 245 0	248.5 249.2 249.9 250.6 251.3		302 0 302 8 303 6 304 5 305 3	365 366 367 368 369
370 371 372 373 374	328 7 329 5 330 4 331 - 3 332 2	172 7 173.2 173 7 174 8 174 7	177 7 178 3 178 8 179 3 179 8	176 B 177 4 177 9 178 4 179 0	170 6 171.1 171 6 173 3 172 7	239 4 240 0 240 7 241.3 242 0	245.7 246.3 247.0 247.7 248.4	252 0 252.7 253 3 254.0 254 7	290 8 291.6 292.4 293.2 294.0	306 1 307 0 307.8 308 6 309.5	370 371 372 373 374
375 376 377 378 379	333 ¹ 334 0 334 9 335 8 336 7	175.3 175 6 176 3 176 8 177.3	180.4 180 9 181.4 182 0 182 5	179.5 180 0 180 6 181 1 181 6	173 2 173 7 174 3 174 8 175-3	242 6 243.3 243 9 244 6 245.2	249 0 249 7 250 4 251 0 251 7	255 4 256 1 256 8 257 5 258.2	294.8 295.6 296.4 297.2 297.9	310,3 311 1 312,0 312.8 313.6	375 376 377 378 379
380 381 382 383 384	337.5 338.4 339.3 340.2 341.1	177-9 178 4 178 9 179 4 180 0	183 0 183 6 184 1 184 6 185 2	182 1 183 2 183 8 184 3	175 9 176.4 176 9 177 5 178.0	245 9 246.6 247 2 247 9 348.5	252 4 253 0 253 7 254 4 255 1	258 8 259 5 260.2 260 9 261.6	198 7 199 5 300 3 301 1 301 9	314.5 315.3 316.1 316 9 317.8	380 381 382 383 384
385 386 387 388 389	342.0 342 9 343 8 344 6 345-5	180.5 181 0 181.5 182 0 182.6	185.7 186 2 186 8 187.3 187 8	184.8 185 4 185 9 186.4 187 0	178.5 179.1 179.6 180.1 180.6	249 2 249 8 250 5 251 1 251 8	255 7 250 4 257 1 257 7 258 4	262.3 263.0 263.6 264.3 265.0	302.7 303.5 304.2 305.0 305.8	318.6 3:9 4 370 3 321.1 321 9	385 386 387 388 389
390 391 392 393 394	346 4 347:3 348:2 349:1 350:0	183 1 183.6 184.1 184.7 185.2	188 4 188 9 189 4 190 0	187 S 168 0 188.6 189 1	181 7 181.7 182.3 182.8 183 3	252 A 253.I 253.7 254.4 255 O	259 I 259.7 260 4 261 I 261 8	265 7 266 4 267 1 267 8 268 5	306.6 307.4 308.2 309.0 309.8	322.8 323 6 324.4 325 2 326.1	390 391 392 393 394

				(W	eights in	milligra	ms.]				
Ĝ.				Invert and St	Sugar acrose.		Lactose.		Mal	tose.	ĝ
Cuprous Oxide (CurO).	Copper (Cu).	Dextrose.	Invert Sugar.	o.4 Gram Total Sugar.	2 Grams Total Sugar.	CuH#Ou.	CuHzOu + 1 HzO.	C13H24O11 + H5O.	CuH#Ou.	Cı:H24O11 + H2O.	Cuprous Oxide (CusO).
395	350.9	185.7	191.0	190.2	183.9	255.7	262.4	269.1	310.6	326.9	395
396	351.8	186.2	191.6	190.7	184.4	256.3	263.1	269.8	311.4	327.7	396
397	352.6	186.8	192.1	191.3	184.9	257.0	263.8	270.5	312.1	328.6	397
398	353.5	187.3	192.7	191.8	185.5	257.7	264.4	271.2	312.9	329.4	398
399	354.4	187.8	193.2	192.3	166.0	258.3	265.1	271.9	313.7	330.2	399
400	335.3	188.4	193.7	192.9	186.5	259.0	265.8	272.6	314.5	331.1	400
401	356.2	188.9	194.3	193.4	187.1	259.6	266.4	273.3	315.3	331.9	401
402	357.1	189.4	194.8	194.0	187.6	260.3	267.1	274.0	316.1	332.7	402
403	358.0	189.9	195.4	194.5	188.1	260.9	267.8	274.6	316.9	333.6	403
404	358.9	190.5	195.9	195.0	188.7	261.6	268.5	275.3	317.7	334.4	404
405 406 407 408 409	359.7 360.6 361.5 362.4 363.3	191.0 191.5 192.1 192.6 193.1	198.4 197.0 197.5 198.1 198.0	195.6 196.1 196.7 197.2 197.7	189.2 189.8 190.3 190.8 191.4	264.2 264.8	269.1 269.8 270.5 271.1 271.8	276.0 276.7 277.4 278.1 278.8	318.5 319.2 320.0 320.8 321.6	335.2 336.0 336.9 337.7 338.5	405 406 407 408 409
410	364.2	193.7	199.1	198.3	191.9	265.5	272.5	279.5	322.4	339·4	410
411	365.1	194.2	199.7	198.8	192.5	266.1	273.1	280.1	323.2	340·2	411
412	366.0	194.7	200.2	199.4	193.0	266.8	273.8	280.8	324.0	341·0	412
413	366.9	195.2	200.8	199.9	193.5	267.4	274.5	281.5	324.8	341·9	413
414	367.7	195.8	201.3	200.5	194.1	268.1	275.2	282.2	325.6	342·7	414
415	368.6	196.3	201.8	201.0	194.6	268.7	275.8	282.9	326.3	343·5	415
416	369.5	196.8	202.4	201.6	195.2	269.4	276.5	283.6	327.1	344·4	416
417	370.4	197.4	202.9	202.1	195.7	270.1	277.2	284.3	327.9	345·2	417
418	371.3	197.9	203.5	202.6	196.2	270.7	277.8	285.0	328.7	346·0	418
419	372.2	198.4	204.0	203.2	196.8	271.4	278.5	285.6	329.5	346·8	419
420	373.1	199.0	204.6	203.7	197.3	272.0	279.2	286.3	330.3	347.7	420
421	374.0	199.5	205.1	204.3	197.9	272.7	279.8	287.0	331.1	348.5	421
422	374.8	200.1	205.7	204.8	198.4	273.3	280.5	287.7	331.9	349.3	422
423	375.7	200.6	206.2	205.4	198.9	274.0	281.2	288.4	332.7	350.2	423
424	376.6	201.1	206.7	205.9	199.5	274.6	281.9	289.1	333.4	351.0	424
425	377 · 5	201.7	207.3	206.5	200.0	275.3	282.5	289.8	334.2	351.8	425
426	378 · 4	202.2	207.8	207.0	200.6	275.9	283.2	290.5	335.0	352.7	426
427	379 · 3	202.8	208.4	207.6	201.1	276.6	283.9	291.1	335.8	353.5	427
428	380 · 2	203.3	208.9	208.1	201.7	277.2	284.5	291.8	336.6	354.3	428
429	381 · 1	203.8	209.5	208.7	202.2	277.9	285.2	292.5	337.4	355.1	429
430	382.0	204.4	210.0	209.2	202.7	278.5	285.9	293.2	338.2	356.0	430
431	382.8	204.9	210.6	209.8	203.3	279.2	280.5	293.9	339.0	356.8	431
432	383.7	205.5	211.1	210.3	203.8	279.8	287.2	294.6	339.7	357.6	432
433	384.6	206.0	211.7	210.9	204.4	280.5	287.9	295.3	340.5	358.5	433
434	385.5	206.5	212.2	211.4	204.9	281.2	288.6	295.9	341.3	359.3	434
435	386.4	207.1	212.8	212.0	205.5	281.8	289.2	296.6	342.1	360.1	435
436	387.3	207.6	213.3	212.5	206.0	282.5	289.9	297.3	342.9	361.0	436
437	388.2	208.2	213.9	213.1	206.6	283.1	290.6	298.0	343.7	361.8	437
438	389.1	208.7	214.4	213.6	207.1	283.8	291.2	298.7	344.5	362.6	438
439	390.0	209.2	215.0	214.2	207.7	284.4	291.9	299.4	345.3	363.4	439
440	390.8	209.8	215.5	214.7	208.2	285.7	292.6	300.1	346.1	364.3	440
441	391.7	210.3	216.1	215.3	208.8	285.7	293.2	300.8	346.8	365.1	441
442	392.6	210.9	216.6	215.8	209.3	286.4	293.9	301.4	347.6	365.9	442
443	393.5	211.4	217.2	216.4	209.9	287.0	294.6	302.1	348.4	366.8	443
444	394.4	212.0	217.8	216.9	210.4	287.7	295.3	302.8	349.2	567.6	444
445	395.3	212.5	218.3	217.5	211.0	288.3	295.9	303.5	350.0	368.4	445
446	396.2	213.1	218.9	218.0	211.5	289.0	296.6	304.2	350.8	369.3	446
447	397.1	213.6	219.4	218.6	212.1	289.6	297.3	304.9	351.6	370.1	447
448	397.9	214.1	220.0	219.1	212.6	290.3	297.9	305.6	352.4	370.9	448
449	398.8	214.7	220.5	219.7	213.2	290.9	298.6	306.3	353.2	371.7	449

				[11	eignts in	mmgra	ns.j		,		
ĝ					Sugar ucrose.		Lactore.		Malt	ose.	Q.
Cuprous Oxide (CusO).	Copper (Cu).	Dextrose.	Invert Sugar.	o.4 Gram Total Sugar.	2 Grams Total Sugar.	CuHzOu.	CisHz:Ou + 1 HzO.	C13H24O11 + H2O.	CuH#Ou.	C13H24O11 + H3O.	Cuprous Oxide (CusO).
450 451 452 453 454	399·7 400.6 401.5 402.4 403.3	215.2 215.8 216.3 216.9 217.4	221.1 221.6 222.2 222.8 223.3	220.2 220.8 221.4 221.9 222.5	213.7 214.3 214.8 215.4 215.9	291.6 292.3 292.9 293.6 294.2	299.3 299.9 300.6 301.3 302.0	306.9 307.6 308.3 309.0	353.9 354.7 355.5 356.3 357.1	372.6 373.4 374.2 375.1 375.9	450 451 452 453 454
455	404.2	218.0	223.9	223.0	216.5	294.9	302.6	310.4	357.9	376.7	455
456	405.1	218.5	224.4	223.6	217.0	295.5	303.3	311.1	358.7	377.6	456
457	405.9	219.1	225.0	224.1	217.6	296.2	304.0	311.8	359.5	378.4	457
458	406.8	219.6	225.5	224.7	218.1	296.8	304.6	312.4	360.3	379.2	458
459	407.7	220.2	220.1	225.3	218.7	297.5	305.3	313.1	361.0	380.0	459
460	408.6	220.7	226.7	225.8	219.2	298.1	306.0	313.8	361.8	380.9	460
461	409.5	221.3	227.2	226.4	219.8	298.8	306.6	314.5	362.6	381.7	461
462	410.4	221.8	227.8	226.9	720.3	299.4	307.3	315.2	363.4	382.5	462
463	411.3	222.4	228.3	227.5	220.9	300.1	308.0	315.9	364.2	383.4	463
464	412.2	222.9	228.9	228.1	221.4	300.7	308.7	316.6	365.0	384.2	464
465	413.0	223.5	229.5	228.6	222.0	301.4	309.3	317.3	365.8	385.0	465
466	413.9	224.0	230.0	229.2	222.5	302.0	310.0	317.9	366.6	385.9	466
467	414.8	224.6	230.6	229.7	223.1	302.7	310.7	318.6	367.3	386.7	467
468	415.7	225.1	231.2	230.3	223.7	303.3	311.3	319.3	368.1	387.5	468
469	416.6	225.7	231.7	230.9	224.2	304.0	312.0	320.0	368.9	388.3	469
470	417.5	226.2	232.3	231.4	224.8	304.7	312.7	320.7	369.7	389.2	470
471	418.4	226.8	232.8	232.0	225.3	305.3	315.3	321.4	370.5	390.0	471
472	419.3	227.4	233.4	232.5	225.9	306.0	314.0	322.1	371.3	390.8	472
473	420.2	227.9	234.0	233.1	226.4	306.6	314.7	322.8	372.1	391.7	473
474	421.0	228.5	234.5	233.7	227.0	307.3	315.4	323.4	372.9	392.5	474
475	421.9	229.0	235.1	234.2	227.6	307.9	316.0	324.1	373.7	393·3	475
476	422.8	229.6	235.7	234.8	228.1	308.6	316.7	324.8	374.4	394·2	476
477	423.7	230.1	236.2	235.4	228.7	309.2	317.4	325.5	375.2	395·0	477
478	424.6	230.7	236.8	235.9	229.2	309.9	318.0	326.2	376.0	395·8	478
479	425.5	231.3	237.4	236.5	229.8	310.5	318.7	326.9	376.8	396·6	479
480	426.4	231.8	237.9	237.1	230.3	311.2	319.4	327.6	377.6	397 · 5	480
481	427.3	232.4	238.5	237.6	230.9	311.8	320.0	328.2	378.4	398 · 3	481
482	428.1	232.9	239.1	238.2	231.5	312.5	320.7	328.9	379.2	399 · 1	482
483	429.0	233.5	239.6	238.8	232.0	313.1	321.4	329.6	380.0	400 · 0	483
484	429.9	234.1	240.2	239.3	232.6	313.8	322.1	330.3	380.7	400 · 8	484
485	430.8	234.6	240.8	239.9	233.2	314.4	322.7	331.0	381.5	401.6	485
486	431.7	235.2	241.4	240.5	233.7	315.1	323.4	331.7	382.3	402.4	486
487	432.6	235.7	241.9	241.0	234.3	315.8	324.1	332.4	383.1	403.3	487
488	433.5	266.3	242.5	241.6	234.8	316.4	324.7	333.1	383.9	404.1	488
489	434.4	236.9	243.1	242.2	235.4	317.1	325.4	333.7	384.7	404.9	489
490	435.3	237.4	243.6	242.7	236.0	317.7	326.1	334.4	385.5	405.8	490

Allihn's Method for the Determination of Dextrose.*—The solutions used are those described on page 501, except that 125 grams of potassium hydroxide are used in place of 50 grams of sodium hydroxide in preparing the alkaline tartrate solution. Place 30 cc. of Fehling's copper solution, 30 cc. of the alkaline tartrate solution, and 60 cc. of water in a beaker and heat to boiling. Add 25 cc. of the sugar solution, which must be so prepared as not to contain more than $I_{i,0}^{C}$ dextrose, and boil over the flame for two minutes. Filter immediately without diluting through a Gooch crucible containing a layer of asbestos fiber, prepared as described on page 504, and wash thoroughly with hot water, using reduced pressure. Transfer the asbestos fiber and the adhering cuprous oxide by means of a glass rod to a beaker and rinse the crucible with about 30 cc. of a boiling mixture of dilute sulphuric and nitric acids containing 65 cc. of sulphuric acid (specific gravity 1.84) and 50 cc. of nitric acid (specific gravity 1.42) per liter. Heat and agitate till the solution is complete, then filter into a scrupulously clean, tared platinum dish of 100-cc. capacity, taking care to wash out all the copper solution from the filter into the dish. Deposit the copper electrolytically in the platinum dish and weigh. Determine the dextrose from Allihn's table. p. 609.

Or, the metallic copper may be calculated by means of the factor 0.7989 from the cupric oxide obtained as in Defren's method (p. 594) and Allihn's table used.

Or, the cuprous oxide as directly obtained by either Allihn's or Defren's method may be washed with alcohol and ether, dried for twenty minutes at 100° C., and weighed, its equivalent in dextrose being ascertained from Allihn's table.

Electrolytic Apparatus.—The author has devised the apparatus shown in Fig. 110 for the electrolytic deposition of copper in sugar analysis and for other work of like nature. A, Fig. 110, is a hard-rubber plate 50 cm. long and 25 cm. wide provided with four insulated metal binding posts, B, each carrying at the top a thumb screw by which a coiled platinum wire electrode, C, may be attached. In front of each post is a copper plate about 4 cm. square covered with thin platinum foil, P, which is bent around the edges of the copper plate and so held in place, the copper plate being screwed to the rubber from beneath. On the square platinum-covered plate is set the platinum evaporating-dish which holds the solution from which the copper is to be deposited, the inside of the dish forming the cathode, while the electrode C, dipping below the surface of the solution, forms the anode. In front of each platinum-covered plate

^{*} Tour. für praktische Chemie, 22 (1880), p. 46.

ALLIHN'S TABLE FOR THE DETERMINATION OF DEXTROSE.*

	ALLUIT		TO LO		IIE DE			011 01		ROSE	·
Milli- grams of	Milli- grams of Cu-	Milli- grams of	Milli- grams of	Milli- grams of Cu-	Milli- grams of	Milli- grams of	Milli- grams of Cu-	Milli- grams of	Milli- grams of	Milli- grams of Cu-	Milli- grams of
Cop- per.	Oxide.	Dex- trose.	Cop- per.	Oxide.	Dex- trose.	Cop- per.	prous Oxide.	Dex- trose.	Cop- per.	prous Oxide.	Dex- trose.
11	12.4	6.6	76	85.6	38.8	141	158.7	71.8	206	231.9	105.8
12	13.5 14.6	7.1	77 78	86.7 87.8	39.3 39.8	I42 I43	159.9	72.3 72.9	207	233.0 234.2	106.3 106.8
14	15.8	7.6 8.1 8.6	79 80	88.9	40.3 40.8	144	162.1	73.4	209	235.3 236.4	107.4
15	16.9			90.1		145	163.2	73.9			107.9
16 17	18.0	9.0	81 82	91.2	41.3 41.8	146	164.4	74.4 74.9	211	237.6 238.7	108.4
17 18	20.3	10.0	83	93.4	42.3	147	165.5	75.5	- 213	239.8	109.5
19 2 0	21.4	10.5	84 85	94.6 95.7	42.8 43.4	149 150	167.7	76.0 76.5	214 215	240.9 242.I	110.0
91	23.6	11.5	86	96.8	43.9	151	170.0	77.0	216	243.2	111.1
22 23	24.8	12.0	87 88	97.9 99.1	44.4	152	171.1	77.5 78.1	217	244.3	111.6
24	25.9 27.0	13.0	89	100.2	45.4	153 154	173.4	78.6	219	245.4 246.6	112.7
25	28.1	13.5	90	101.3	45.9	155	174.5	79.1	220	247.7	113.2
26	29.3	14.0	91	102.4	46.4 46.9	156	175.6	79.6 80.1	221	248.7	113.7
27 28	30.4 31.5	14.5 15.0	92 93	104.7	47.4	157	177.9	80.7	223	249.9 251.0	114.3
29 30	32.7	15.5	94 95	105.8	47.9 48.4	159 160	179.0	81.2 81.7	224	252.4 253.3	115.3
			1			_		1		!	1
31 32	34.9 36.0	16.5	96 97	108.1	48.9 49.4	161	181.3	82.2 82.7	226	254.4 255.6	116.4
33	37.2	17.5	98	110.3	49.9	163	183.5	83.3 83.8	228	256.7	117.4
34 35	38.3	18.0	100	111.5	50.4	164	184.6 185.8	84.3	229 230	257.8 258.9	118.0
36	40.5	18.9	101	113.7	51.4	166	186.9	84.8	231	260.I	119.0
37 38	41.7	19.4	102	114.8	51.9	167	188.0	85.3	232	261.2	119.6
38 39	42.8	19.9	103	116.0	52.4 52.9	168	189.1	85.9 86.4	233 234	262.3 263.4	120.1
40	45.0	20.9	105	118.2	53-5	170	191.4	86.9	235	264.6	121.2
41	46.2	21.4	106	119.3	54.0	171	192.5	87.4	236	265.7 266.8	121.7
42 43	47.3 48.4	21.9	107	120.5	54·5 55·0	172	193.6	87.9 88.5	237	268.0	122.3
44 45	49.5	22.9	109	122.7	55.5 56.0	174 175	195.9	89.0 89.5	230 240	269. I 270. 2	123.4
			ll .	Ì	1 .	<u> </u>					
46 47	51.8	23.9 24.4	111	125.0	56.5 57.0	176	198.1	90.0	24I 242	271.3 272.5	124.4
47 48	54.0	24.9	113	127.2	57·5 58.0	178	200.4	91.1	243	273.6	125.5
49 50	55.2 56.3	25.4 25.9	114	120.3	58.6	179	201.5	92.1	244 245	274.7 275.8	126.6
51	57.4	26.4	116	130.6	59.1	181	203.8	92.6	246	277.0	127.1
52	58.5 59.7	26.9 27.4	117	131.7	59.6 60.1	182	204.9	93.1 93.7	247 248	278.1 279.2	127.6
53 54	60.8	27.9	119	134.0	60.6	184	207.1	94.2	249	280.3	128.7
55	61.9	28.4	120	135.1	61.1	185	208.3	94.7	250	281.5	129.2
56	63.0	28.8	121	136.2	61.6 62.1	186 187	209.4	95.2	251	282.6	129.7
57 58	64.2 65.3 66.4	29.3	122 123	137.4 138.5	62.6	188	210.5	95.7 96.3	252 253	283.7 284.8 286.0	130.3
59 60	66.4	30.3 30.8	124	139.6	63.1 63.7	189	212.8	96.8 97.3	254 255	286.0 287.1	131.4
61	68.7		126	141.0	64.2	191	I	97.8	256	288.2	1
62	69.8	31.3 31.8	127	143.0	64.7	192	215.0 216.2	08.4	257	289.3	132.4
63 64	70.9 72.1	32.3 32.8	128	144.1	65.2	193	217.3	98.9 90.4	258 259	290.5 291.6	133.5 134.1
65	73.2	33.3	130	145.2 146.4	66.2	195	219.5	100.0	260	292.7	134.6
66	74.3	33.8	131	147.5	66.7	196	220.7	100.5	261	293.8	135.1
67 68	75.4 76.6	34.3 34.8	132	148.6	67.2	197	221.8	101.0	262 263	205.0 206.1	135.7 136.2
69	77.7	35.3 35.8	134	150.0	68.2	199	224.0	102.0	264	207.2	136.8
70	78.8	35.8	135	152.0	68.8	200	225.2	102.6	265	298.3	137.3
71	79.9 81.1	36.3 36.8	136	153.1	69.3 69.8	201	226.3	103.1 103.7	266 267	299.5 300.6	137.8
72 73	82.2	37.3 37.8	137	154.2 155.4 156.5	70.3	203	227.4	104.2	268	301.7	138.9
74 75	83.3 84.4	37.8 38.3	139	156.5 157.6	70.8	204	229.7 230.8	104.7	269	302.8	139.5
/3		33.3	- 40	-37.0	,	1 203		3.3	1 3/0	3-4.0	

^{*} U. S. Dept. of Agric.. Bur. of Chem.. Bul. 65.p 143

ALLIHN'S TABLE FOR THE DETERMINATION OF DEXTROSE—(Continued).

										(00,0	· · · · · · · · · · · · · · · · · · ·
Milli-	Milli-	Milli-	Milli-	Milli-	MiDi-	Milli-	Milli-	Milli-	Milli-	Milli-	Milli-
grams	grams	grams	grams	grams	grams	grams	grams		grams	grams	grams
of	of Cu-	of	of	of Cu-	of	of	of Cu-	of	of	of Cu-	of
Cop-	prous	Dex-	Cop-	prous	Dex-	Cop-	prous		Cop-	prous	Dex-
per.	Oxide.	trose.	per.	Oxide.	trose.	per.	Oxide.	trose.	per.	Oxide.	trose.
<u> </u>			! 						: -		 -
271	305 . I	140.6	321	361.4	168.1 168.6	371	417.7	196.3	421	474.0	225.1
272 273	306.2	141.1	322	362.5 363.7	160.2	372 373	418.8	196.8	422	475.6 476.2	225.7 226.3
274	307.3 308.5	142.3	324	364.8	160.7	374	421.1	108.0	424	477-4	226.0
275	300.5	142.8	325	365.9	170.3	375	422.2	198.6	425	478.5	227.5
-//3	309.0		3-3		.,	1	4		7-3	4/0.3	,.,
276	310.7	143.3	326	367.0	170.9	376	423.3	199.1	426	479.6	228.0
277	311.9	143.9	327	368.2	171.4	377	424.5	199.7	427	480.7	228.6
278	313.0	144.4	328	369.3	172.0	378	425.6	200.3	428	481.9	229.2
279 280	314.1	145.0	329	370.4	172.5	379 380	426.7	200.8	429	483.0 484.1	229.8 230.4
260	315.2	145.5	330	371.5	173.1	380	427.6	201.4	430	404.1	230.4
281	316.4	146.1	331	372.7	173.7	38 r	429.0	202.0	431	485.3	231.0
282	317.5	146.6	332	373.8	174.2	382	430. I	202.5	432	486.4	231.6
283	318.6	147.2	333	374.9	174.8	383	431.2	203.I	433	487.5	232.2
284	319.7	147.7	334	376.0	175.3	384		203.7	434	488.6	232.8
285	320.9	148.3	335	377.2	175.9	385	433.5	204.3	435	489.7	233.4
286	322.0	148.8	336	378.3	176.5	386	434.6	204.8	436	490.9	233.9
287	323.I	149.4	337	379.4	177.0	387	435.7	205.4	437	492.0	234.5
288	324.2	149.9	338	380.5	177.6	388	436.8	206.0	438	493.I	235.I
289	325.4	150.5	339	381.7	178.1	389	438.0	206.5	439	494 - 3	235.7
290	326.5	151.0	340	382.8	178.7	390	439.1	207.1	440	495 - 4	236.3
201	327.4	151.6	341	383.9	179.3	301	440.2	207.7	441	496.5	236.9
202	328.7	152.1	342	385.0	179.8	392	441.3	208.3	442	497.6	237.5
293	329.9	152.7	343	386.2	180.4	393	442.4	208.8	443	498.8	238.1
294	331.0	153.2	344	387.3	180.9	394	443.6	209.4	444	499.9	238.7
295	332.1	153.8	345	388.4	181.5	395	444 - 7	210.0	445	501.0	239.3
296	333.3	154.3	346	389.6	182.1	396	445.9	210.6	446	502.1	239.8
297	334.4	154.9	347	390.7	182.6	397	447.0	211.2	447	503.2	240.4
298	335 - 5	155.4	348	391.8	183.2	398	448.I	211.7	448	504.4	241.0
299	336.6	156.0	349	392.9	183.7	399	440.2	212.3	449	505.5	241.6
300	337.8	156.5	350	394.0	184.3	400	450.3	212.9	450	506.6	242.2
301	338.0	157.1	35T	305.2	184.0	401	451.5	213.5	451	507.8	242.8
302	340.0	157.6	352	396.3	185.4	402	452.6	214.1	452	508.9	243.4
303	341.1	158.2	353	397 - 4	186.0	403	453.7	214.6	453	510.0	244.0
304	342.3	158.7	354	398.6	186.6	404	454.8	215.2	454	511.1	244.6
305	343-4	159.3	355	399 - 7	187.2	405	456.0	215.8	455	512.3	245.2
306	344 - 5	150.8	356	400.8	187.7	406	457.I	216.4	456	513.4	245.7
307	345.6	160.4	357	401.9	188.3	407	458.2	217.0	457	514.5	246.3
308	346.8	160.9	358	403.I	188.9	408	459-4	217.5	458	515.6	246.9
300	347.9	161.5	359	404.2	189.4	409	460.5	218.1	459	516.8	247.5
310	349.0	162.0	360	405.3	190.0	410	461.6	218.7	460	517.9	248. I
311	350.1	162.6	361	406.4	100.6	411	462.7	219.3	461	510.0	248.7
312	351.3	163.1	362	407.6	191.1	412	463.8	219.9	462	520.I	249.3
313	352.4	163.7	363	408.7	191.7	413	465.0	220.4	463	521.3	249.9
314	353.5	164.2	364	409.8	192.3	414	466.1	221.0			
315	354.6	164.8	365	410.9	192.9	415	467.2	221.6			
316	355.8	165.3	366	412.1	193.4	416	468.4	222.2			
317	356.9	165.9	367	413.2	194.0	417	469.5	222.8			1
318	358.0	166.4	368	414.3	194.6	418	470.6	223.3	ll l		
319	359.1	167.0	369	415.4	195.1	419	471.8	223.9			
320	360.3	167.5	370	416.6	195.7	420	472.9	224.5			
1	-							1	ll .		

is a switch, S, and at either end of the hard-rubber plate is a binding post, R, for connection with the electric current. The wiring, which is on the under side of the rubber plate, is best illustrated by the diagram in Fig. 110.

Four determinations may be carried on simultaneously in four platinum dishes, if desired, the wiring and the switches being so arranged that beginning at one end of the plate either the first dish or the first

Fig. 110.—Four Pan Electrolytic Apparatus, shown (above) with Glass-covered Top Partially Removed, and (below) in Diagram.

two or the first three may be thrown in or out of circuit at will without interrupting the current through the remaining dishes. A cover with wooden sides and glass top fits closely over the whole apparatus as a protection from dust, but may be easily lifted off to manipulate the dishes when desired. The sides of the cover are perforated to permit the escape of the gas formed during the electrolysis.

The ordinary street current is used when available, and the strength of the current may be varied within wide limits by means of a number of 16 or 32 candle-power lamps, K, coupled in multiple, and a rheostat, L, consisting of a vertical glass tube sealed at the bottom, containing a column of dilute acid, the resistance being changed by varying the length of the acid column contained between the two platinum terminals immersed therein, one of which is movable. A gravity battery of four cells may be employed if the laboratory is not equipped with electric lights.

In using this apparatus for determining copper, as in sugar work, the plating process should go on till all the copper is deposited, requiring several hours or over night with a current strength of about 0.25 ampere. Before stopping the process, the absence of copper in the solution should be proved by removing a few drops with a pipette, adding first ammonia, then acetic acid, and testing with ferrocyanide of potassium. If no brown coloration is produced, all the copper has been plated out. Throw the dish out of circuit by means of the switch, pour out the acid solution quickly before it has a chance to dissolve any of the copper, wash the dish first with water and then with alcohol, dry, and weigh.

The copper may be removed from the platinum dish by strong nitric acid.

Determination of Sucrose by Fehling's Solution.*—If a polariscope is not available, cane sugar can be determined as follows: First determine the percentage of invert sugar present in the sample by one of the Fehling methods already described. Then dissolve I gram of the sugar in about 100 cc. of water in a 500-cc. graduated flask, add 3 cc. of concentrated hydrochloric acid and invert by heating in water to 68° and cooling in the regular manner. Neutralize with sodium hydroxide or sodium carbonate, and make up to the mark with water. Determine the per cent of total reducing sugar as invert sugar either by the volumetric or gravimetric Fehling process. Subtract the invert sugar found present in the sugar by direct determination from the total found present after inversion, and

^{*} Tucker, Manual of Sugar Analysis, p. 182.

the remainder is the invert sugar due to cane sugar. This figure multiplied by 0.05 gives the percentage of cane sugar.

For the determination of sucrose by the gravimetric Fehling process on the inverted sample, multiply the cupric oxide (CuO) by the factor 0.4307, or the copper (Cu) by the factor 0.5394.

ANALYSIS OF MOLASSES AND SYRUPS.

First insure a perfectly homogeneous sample by stirring with a rod to evenly distribute any separated sugar.

Determination of Total Solids.—(1) Asbestos Method.—Weigh 20 grams into a 100-cc. graduated flask, dissolve in water, and make up to the mark. Insure a uniform solution by shaking. Measure 10 cc. of this solution into a tared platinum dish containing about 5 grams of freshly ignited, finely divided asbestos fiber, and dry to constant weight at 70° in vacuo, or in a McGill oven (see p. 586).

- (2) Sand Method.*—Place about 15 grams of ignited quartz sand and a stirring rod in a flat-bottom metal dish and weigh. Add 2 to 4 grams of the material and sufficient moisture to permit thorough mixing. Dry on a water bath with stirring and finally in a water oven until the loss in weight in one hour is not more than 3 mg. At least 8 hours' heating is usually required.
- (3) By Calculation from Refractive Index.—Determine the refractive index by means of the Abbé refractometer (p. 108), and calculate the total solids, using Geerligs's tables (p. 615).

This method is more accurate and convenient than the specific gravity method and employs a smaller quantity of material. The investigations of Stolle† and of Tolman and Smith‡ have shown that sucrose, maltose, dextrose, levulose and lactose all have practically the same refractive index. Dextrin has a somewhat higher refractive index, nevertheless the solids of commercial glucose do not give a reading appreciably higher than the sugars named.

A. H. Bryan, has compared this method with the method of drying at 70° in vacuo, with the following results:

^{*} U. S. Dept. Agric., Bur. of Chem., Bul. 107 (rev.), p. 65.

[†] Zeits. deutsch. Zucker-Ind., 1901, pp. 335, 469.

[‡] Jour. Am. Chem. Soc., 28, 1906, p. 1476.

[§] Ibid., 30, 1908, p. 1443.

Material.	Number of Samples.	Difference compared with the Gravimetric Method.
Maple syrup	13	-1.34 to $+0.72$
Cane table syrup	10	-0.79 " +0.62
Cane molasses	17	-1.53 " +0.59
Beet molasses	15	-1.83 " -0.07
Honey	24	-2.52 " +0.91
Glucose	2	-0.27 " +0.27

(4) By Calculation from Specific Gravity.—Weigh 25 grams of the sample into a 100-cc. graduated flask, dissolve in water, and make up to the mark. Determine the specific gravity of the diluted solution by means of a pycnometer or Westphal balance. Ascertain from the table on pp. 617-620 the percentage by weight of solids (sugar) corresponding to the specific gravity of the diluted solution, and calculate the total solids in the original sample by the formula

Solids in original sample = 4DS,

D being the specific gravity of the diluted solution and S the per cent of solids in the diluted solution.

Determination of Ash.—Weigh from 5 to 10 grams of the sample into a tared platinum dish, evaporate to dryness on the water-bath, and proceed as directed for ash of sugar (p. 586).

Polarization and Determination of Sucrose.—Molasses and golden syrup require the application of clarifying reagents before a sufficiently clear solution can be obtained for reading on the polariscope. Even then it is not possible nor is it necessary to get a water-white solution, so that in this class of products greater accuracy can usually be attained by polarizing in a 100-mm. tube (half the standard length) and multiplying the reading by 2. The clarifier best adapted as a rule for molasses and golden syrup is subacetate of lead.*

The Process.—The normal weight, 26.048 grams, of the molasses or syrup is dissolved in water in a 100-cc. flask, and in the case of molasses and "golden," or "drip" syrup, sufficient subacetate of lead solution is added to precipitate the coloring matter. From 5 to 10 cc. of the clarifier usually suffice. The flask is then filled to the mark with water and the contents shaken thoroughly and filtered. If on account of air bubbles it is difficult to make up to the mark, the bubbles may usually be dispelled by a drop of ether. With maple syrup no clarifier is, as a rule, necessary, though sometimes alumina cream is helpful. With a very

^{*} Alumina cream, p. 587, and bone black, or animal char, are also useful.

GEERLIGS'S TABLE FOR DRY SUBSTANCE IN SUGAR-HOUSE PRODUCTS
BY THE ABBE REFRACTOMETER, AT 28° C.*

Refrac- tive Index.	Per Cent Dry Sub- stance.	Decimals to b		Refrac- tive Index.	Per Cent Dry Sub- stance.	Decimals to be Added for Fractional Readings †			
1.3335	ı	0.0001=0.05	0.0010=0.75	1.4083	45	0.0004=0.2 0.0015=0.75			
1.3349	2	0.0002=0.1	0.0011=0.8	1.4104	46	0.0005=0.250.0016=0.8			
1.3364	3	0.0003=0.2	0.0012=0.8	1.4124	47	0.0006=0.3 0.0017=0.85			
1.3379	4	0.0004=0.25	0.0013=0.85	1.4145	48	0.0007=0.35 0.0018=0.0			
1.3394		0.0005=0.3	0.0014=0.0	1.4166	49	0.0008=0.4 0.0019=0.05			
1.3409	5 6	0.0006=0.4	0.0015=1.0	1.4186	50	0.0009=0.450.0020=1.0			
1.3424	7	0.0007=0.5		1.4207	51	0.0010=0.5 0.0021=1.0			
1.3439	8	0.0008=0.6		1.4228	52	0.0011=0.55			
1.3454	9	0.0009=0.7		1.4219		33			
1.3469	10			1.4270					
1.3484	11	0.0001=0.05		1.4292	55	0.0001=0.050.0013=0.55			
1.3500	12	0.0002=0.I		1.4314	56	0.0002=0.1 0.0014=0.6			
1.3516	13	0.0003=0.2		1.4337	57	0.0003=0.1 0.0015=0.65			
1.3530	14	0.0004=0.25		1.4359	58	0.0004=0.15 0.0016=0.7			
1.3540	15	0.0005=0.3		1.4382	59	0.0005=0.2 0.0017=0.75			
1.3562	16	0.0006=0.4		1.4405	60	0.0006=0.25 0.0018=0.8			
1.3578	17	0.0007=0.45		1.4428	61	0.0007=0.3 0.0019=0.85			
1.3594	18	0.0008=0.5		1.4451	62	0.0008=0.35 0.0020=0.9			
1.3611	19	0.0009=0.6		1.4474	63	0.0009=0.4 0.0021=0.9			
1.3627	20 21	0.0010=0.65		1.4497	64	0.0010=0.45 0.0022=0.95			
1.3644 1.3661	21	0.0011=0.7		1.4520	65	0.0011=0.5 0.0023=1.0			
1.3678	1	0.0012=0.75		1.4543	66	0.0012=0.5 0.0024=1.0			
1.3695	23	0.0013=0.8		1.4567	67 68				
1.3712	25	0.0014=0.85 0.0015=0.0		1.4591					
1.3729	26	0.0015=0.95		1.4615	69				
31-9	••	0.93		1.4663	70 71				
				1.4687	72				
1.3746	27	0.0001=0.05	0.0012=0.6	1.400/	/-				
1.3764	28	0.0002=0.I	0.0013=0.65						
1.3782	29	0.0003=0.15	0.0014=0.7	1.4711	73	0.0001=0.0 0.0015=0.55			
1.3800	30	0.0004=0.2	0.0015=0.75	1.4736	74	0.0002=0.050.0016=0.6			
1.3818	31	0.0005=0.25	0.0016=0.8	1.4761	75	0.0003=0.1 0.0017=0.66			
1.3836	32	0.0006=0.3	0.0017 = 0.85	1.4786	76	0.0004=0.150.0018=0.66			
1.3854	33	0.0007=0.35	0.0018=0.9	1.4811	77	0.0005=0.2 0.0010=0.7			
1.3872	34	0.0008=0.45	0.0019=0.95	1.4836	78	0.0006=0.2 0.0020=0.75			
1.3890	35	0.0009=0.4	0.0020=1.0	1.4862	79	0.0007=0.25 0.0021=0.8			
1.3909	36	0.0010=0.5	0.0021=1.0	1.4888	80	0.0008=0.3 0.0022=0.8			
1.3928	37	0.0011=0.55		1.4914	81	0.0009=0.35 0.0023=0.85			
1.3947	38			1.4940	82	0.0010=0.35 0.0024=0.9			
1.3966	39			1.4966	83	0.0011=0.4 0.0025=0.9			
1.3984	40			1.4992	84	0.0012=0.45 0.0026=0.95			
1.4003	41			1.5019	85	0.0013=0.5 0.0027=1.0			
				1.5046	86	0 0014=0.5 0.0028=1.0			
1.4023	42	0.0001=0.05	0.0012=0.6	1.5073	87 88				
1.4043	43	0.0002=0.1	0.0013=0.65	1.5127	89				

*Intern. Sugar Jour., 10, pp. 69-70.
† Find in the table the refractive index which is next lower than the reading actually made and note the corresponding whole number for the per cent of dry substance. Subtract the refractive index obtained from the table from the observed reading; the decimal corresponding to this difference, as given in the column so marked, is added to the whole per cent of dry substance as first obtained.

TEMPERATURE	CORRECTIONS	FOR 1	USE	WITH	GEERLIGS'S	TABLE.

Tempera-		Dry Substance.														
ture of the Prisms in	0	5	10	15	20	25	30	40	50	60	70	80	90			
°C.		. Subtract—														
20	0.53	0.54	0.55	0.56	0.57	0.58	0.60	0.62	0.64	0.62	0.61	0.60	0.58			
21	.46	-47	.48	-49	.50	.51	-52	-54	.56	- 54	-53	-52	-50			
22	.40	.41	-42	-42	-43	-44	-45	-47	.48	-47	.46	-45	-44			
23	-33	-33	-34	-35	.36	-37	.38	-39	.40	-39	.38	.38	.38			
24	.26	. 26	.27	. 28	. 28	.29	.30	.31	.32	.31	-31	.30	.30			
25	.20	.20	.21	.21	.22	.22	.23	:23	.24	-23	. 23	.23	.22			
26	.12	.12	.13	.14	.14	.15	.15	.16	.16	.16	-15	.15	-14			
27	.07	-07	-07	-07	.07	-07	.08	.08	.08	.08	.08	.08	.07			
							Add-	_								
29	0.07	0.07	0.07	0.07	0.07	0.07	0.08	0.08	0.08	0.08	0.08	0.08	0.07			
30	.12	.12	.13	.14	-14	-14	.15	.15	.16	.16	.16	.15	.14			
31	.20	.20	.21	.21	.22	.22	.23	.23	.24	-23	-23	.23	. 22			
32	.26	.26	.27	.28	. 28	.29	.30	-31	-32	-31	.31	-30	-30			
3 3	-33	-33	-34	-35	.36	-37	-38	-39	.40	-39	.38	.38	.38			
34	.40	-41	.42	-42	-43	-44	-45	-47	-48	-47	-46	-45	-44			
35	.46	-47	.48	1 -49	- 50	.51	-52	-54	1.56	- 54	- 53	.52	-50			

dark-colored molasses 20 to 30 cc. of lead subacetate are required for clarification and in extreme cases (though rarely with the grades of molasses used as food) it is necessary, after the ordinary filtration, to pass through from 5 to 6 grams of powdered, dried bone charcoal.*

An excess of subacetate of lead should be avoided on account of the possibility of the filtrate becoming turbid through the formation of lead carbonate by exposure to the air. A drop of acetic acid will nearly always clear the solution, if the turbidity is due to carbonate. If cloudiness in the filtrate persists, weigh out a fresh portion of the sample, dilute, and add first the lead subacetate solution, and afterwards enough of a strong solution of sodium sulphate or common salt to precipitate the excess of lead; then fill to the mark and filter. Polarize, and conduct the inversion as directed on p. 588, using, however, a 100-mm. tube, and multiplying the reading by 2, both direct and invert.† Use Clerget's formula for calculation of the sucrose.

For medium- or light-colored grades of molasses, which yield but a small precipitate with lead subacetate, the above method of simple polarization, both direct and invert, gives results sufficiently accurate for ordinary work. For dark-colored, or "black-strap" molasses, or wherever

^{*} The treatment with bone char should be used only as a last resort, as, on account of slight absorption of sugar, observed readings are from 0.4° to to 0.5° too low.

[†] The short tube (100 mm.) is preferred for polarizing molasses, not only on account of the more or less deep color of the clarified solution, but also because a molasses sample con. taining considerable commercial glucose would not read within the scale limits, if the 200-mmtube were employed.

RELATION OF BRIX, SPECIFIC GRAVITY, AND BAUMÉ.

Per Cent of	Specific Gravity.	Degree Baumé.	Per Cent of	Specific	Degree Baumé.	Per Cent of	Specific Gravity.	Degree Baumé.	Per Cent of	Specific	Degree Baumé.
Sugar.	Giavity.	Qa	Sugar.	Gravity.	Qm	Sugar.	Giavity.	Q _m	Sugar.	Gravity.	A A
0.1 0.2 0.3 0.4 0.5	1.0003 1.0007 1.0011 1.0015 1.0019	0.06 0.11 0.17 0.22 0.28	6.6 6.7 6.8 6.9 7.0	1.0261 1.0265 1.0269 1.0273 1.0277	3.7 3.7 3.8 3.8 3.9	13.1 13.2 13.3 13.4 13.5	1.0531 1.0536 1.0540 1.0544 1.0548	7·3 7·3 7·4 7·4 7·5	19.6 19.7 19.8 19.9 20.0	1.0815 1.0819 1.0824 1.0828 1.0832	10.85 10.9 11.0 11.0
0.6 0.7 0.8 0.9	1.0023 1.0027 1.0031 1.0034 1.0038	0.33 0.39 0.44 0.5 0.55	7.1 7.2 7.3 7.4 7.5	1.0281 1.0286 1.0290 1.0294 1.0298	3.9 4.0 4.1 4.1 4.2	13.6 13.7 13.8 13.9 14.0	1.0553 1.0557 1.0561 1.0566 1.0570	7·5 7·6 7·65 7·7 7·8	20.1 20.2 20.3 20.4 20.5	1.0837 1.0841 1.0846 1.0850 1.0855	11.1 11.2 11.2 11.3
1.1 1.2 1.3 1.4 1.5	1.0042 1.0046 1.0050 1.0054 1.0058	o.6 o.7 o.7 o.8 o.8	7.6 7.7 7.8 7.9 8.0	1.0302 1.0306 1.0310 1.0314 1.0318	4.2 4.3 4.3 4.4 4.4	14.1 14.2 14.3 14.4	1.0574 1.0578 1.0583 1.0587	7.8 7.9 7.9 8.0 8.0	20.6 20.7 20.8 20.9 21.0	1.0859 1.0864 1.0868 1.0873	11.4 11.45 11.5 11.6
1.6 1.7 1.8 1.9	1.0062 1.0066 1.0070 1.0074 1.0077	0.9 0.9 1.0 1.05	8.1 8.2 8.3 8.4 8.5	1.0322 1.0327 1.0331 1.0335 1.0339	4·5 4·55 4·6 4·7 4·7	14.6 14.7 14.8 14.9	1.0506 1.0600 1.0604 1.0609 1.0613	8.1 8.15 8.2 8.3 8.3	21.1 21.2 21.3 21.4 21.5	1.0882 1.0886 1.0891 1.0895	11.7 11.7 11.8 11.8
2.1 2.2 2.3 2.4 2.5	1.0081 1.0085 1.0089 1.0093 1.0097	1.2 1.2 1.3 1.3	8.6 8.7 8.8 8.9 9.0	1.0343 1.0347 1.0351 1.0355 1.0359	4.8 4.8 4.9 4.9 5.0	15.1 15.2 15.3 15.4 15.5	1.0617 1.0621 1.0626 1.0630 1.0634	8.4 8.4 8.5 8.5 8.6	21.6 21.7 21.8 21.9 22.0	1.0904 1.0909 1.0914 1.0918 1.0923	11.95 12.0 12.05 12.1 12.2
2.6 2.7 2.8 2.9 3.0	1.0101 1.0105 1.0109 1.0113 1.0117	1.4 1.5 1.55 1.6	9.1 9.2 9.3 9.4 9.5	1.0364 1.0368 1.0372 1.0376 1.0380	5.05 5.1 5.2 5.2 5.3	15.6 15.7 15.8 15.9 16.0	1.0639 1.0643 1.0647 1.0652 1.0656	8.65 8.7 8.8 8.8 8.9	22.1 22.2 22.3 22.4 22.5	1.0927 1.0932 1.0936 1.0941 1.0945	12.2 12.3 12.3 12.4 12.4
3.1 3.2 3.3 3.4 3.5	1.0121 1.0125 1.0129 1.0133 1.0137	1.7 1.8 1.8 1.9	9.6 9.7 9.8 9.9	1.0384 1.0388 1.0393 1.0397 1.0401	5·3 5·4 5·4 5·5 5·5	16.1 16.2 16.3 16.4 16.5	1.0660 1.0665 1.0669 1.0674 1.0678	8.9 9.0 9.0 9.1 9.1	22.6 22.7 22.8 22.9 23.0	1.0950 1.0954 1.0959 1.0964 1.0968	12.5 12.55 12.6 12.7 12.7
3.6 3.7 3.8 3.9	1.0141 1.0145 1.0149 1.0153 1.0157	2.0 2.0 2.1 2.2 2.2	10.1 10.2 10.3 10.4 10.5	1.0405 1.0409 1.0413 1.0418 1.0422	5.6 5.7 5.7 5.8 5.8	16.6 16.7 16.8 16.9	1.0682 1.0687 1.0691 1.0695 1.0700	9.2 9.25 9.3 9.4 9.4	23.1 23.2 23.3 23.4 23.5	1.0973 1.0977 1.0982 1.0986 1.0991	12.8 12.8 12.9 12.9
4.1 4.2 4.3 4.4 4.5	1.0161 1.0165 1.0169 1.0173 1.0177	2.3 2.3 2.4 2.4 2.5	10.6 10.7 10.8 10.9	1.0426 1.0430 1.0434 1.0430 1.0443	5.9 5.9 6.0 6.05 6.1	17.1 17.2 17.3 17.4	1.0704 1.0709 1.0713 1.0717 1.0722	9.5 9.5 9.6 9.6 9.7	23.6 23.7 23.8 23.9 24.0	1.0996 1.1000 1.1005 1.1009 1.1014	13.0 13.1 13.15 13.2
4.6 4.7 4.8 4.9 5.0	1.0181 1.0185 1.0189 1.0193 1.0197	2.6 2.6 2.7 2.7 2.8	11.1 11.2 11.3 11.4 11.5	1.0447 1.0451 1.0455 1.0459 1.0464	6.2 6.2 6.3 6.3	17.6 17.7 17.8 17.9 18.0	1.0726 1.0730 1.0735 1.0739 1.0744	9.75 9.8 9.9 9.9	24.1 24.2 24.3 24.4 24.5	1.1019 1.1023 1.1028 1.1032 1.1037	13.3 13.4 13.4 13.5 13.5
5.1 5.2 5.3 5.4 5.5	t.0201 t.0205 t.0209 t.0213	2.8 2.9 2.9 3.0 3.0	11.6 11.7 11.8 11.9 12.0	1.0468 1.0472 1.0476 1.0481 1.0485	6.4 6.5 6.55 6.6 6.7	18.1 18.2 18.3 18.4 18.5	1.0748 1.0753 1.0757 1.0761 1.0766	10.0 10.1 10.1 10.2 10.2	24.6 24.7 24.8 24.9 25.0	1.1042 1.1046 1.1051 1.1056 1.1060	13.6 13.7 13.75 13.8
5.6 5.7 5.8 5.9 6.0	1.0221 1.0225 1.0229 1.0233 1.0237	3.1 3.2 3.2 3.3 3.3	12.1 12.2 12.3 12.4 12.5	1.0489 1.0493 1.0497 1.0502 1.0506	6.7 6.8 6.8 6.9 6.9	18.6 18.7 18.8 18.9	1.0770 1.0775 1.0779 1.0783 1.0788	10.3 10.35 10.4 10.5	25.1 25.2 25.3 25.4 25.5	1.1065 1.1070 1.1074 1.1079 1.1083	13.9 13.9 14.0 14.0
6.1 6.2 6.3 6.4 6.5	1.0241 1.0245 1.0249 1.0253 1.0257	3.4 3.5 3.6 3.6	12.6 12.7 12.8 12.9 13.0	1.0510 1.0514 1.0519 1.0523 1.0527	7.0 7.05 7.1 7.2 7.2	19.1 19.2 19.3 19.4 19.5	1.0792 1.0797 1.0801 1.0806 1.0810	10.6 10.6 10.7 10.7 10.8	25.6 25.7 25.8 25.9 26.0	1.1088 1.1093 1.1097 1.1102 1.1107	14.1 14.2 14.2 14.3

RELATION OF BRIX, SPECIFIC GRAVITY, AND BAUME—(Continued).

	. (انه		<u>-</u>	ye: 1		 -	II		1	
Per Cent of Sugar.	Specific Gravity.	Degree Baume	Per Cent of Sugar.	Specific Gravity.	Degree Baume	Per Cent of Sugar.	Specific Gravity.	Degree Baume	Per Cent of Sugar.	Specific Gravity.	Degree Baumé.
26.1 26.2 26.3 26.4 26.5	1.1111 1.1116 1.1121 1.1125 1.1130	14.4 14.5 14.5 14.6 14.6	32.6 32.7 32.8 32.9 33.0	1.1427	17.9 18.0 18.0 18.1 18.15	39.1 39.2 39.3 39.4 39.5	1.1748 1.1753 1.1758 1.1763 1.1768	21.4 21.5 21.5 21.6 21.6	45.6 45.7 45.8 45.9 46.0	1.2088 1.2093 1.2099 1.2104 1.2110	24.9 24.9 25.0 25.0 25.1
26.6 26.7 26.8 26.9 27.0	1.1135 1.1140 1.1144 1.1149 1.1154	14.7 14.7 14.8 14.8	33.1 33.2 33.3 33.4 33.5	1.1447 1.1452 1.1457 1.1462 1.1466	18.2 18.25 18.3 18.4 18.4	39.6 39.7 39.8 39.9 40.0	1.1773 1.1778 1.1784 1.1789 1.1794	21.7 21.7 21.8 21.85 21.9	46.1 46.2 46.3 46.4 46.5	1.2115 1.2120 1.2126 1.2131 1.2136	25.1 25.2 25.2 25.3 25.35
27.1 27.2 27.3 27.4 27.5	1.1158 1.1163 1.1168 1.1172 1.1177	14.9 15.0 15.1 15.1 15.2	33.6 33.7 33.8 33.9 34.0	1.1471 1.1476 1.1481 1.1486 1.1491	18.5 18.5 18.6 18.6 18.7	40.1 40.2 40.3 40.4 40.5	1.1799 1.1804 1.1809 1.1815 1.1820	22.0 22.0 22.1 22.1 22.2	46.6 46.7 46.8 46.9 47.0	1.2142 1.2147 1.2153 1.2158 1.2163	25.4 25.43 25.5 25.6 25.6
27.6 27.7 27.8 27.9 28.0	1.1182 1.1187 1.1191 1.1196 1.1201	15.2 15.3 15.3 15.4 15.4	34.1 34.2 34.3 34.4 34.5	1.1496 1.1501 1.1506 1.1511 1.1516	18.7 18.8 18.85 18.9 18.95	40.6 40.7 40.8 40.9 41.0	1.1825 1.1830 1.1835 1.1840 1.1846	22.2 22.3 22.3 22.4 22.4	47.1 47.2 47.3 47.4 47.5	1.2169 1.2174 1.2180 1.2185 1.2191	25.7 25.7 25.8 25.8 25.8
28.1 28.2 28.3 28.4 28.5	1.1206 1.1210 1.1215 1.1220 1.1225	15.5 15.55 15.6 15.7 15.7	34.6 34.7 34.8 34.9 35.0	1.1521 1.1526 1.1531 1.1536 1.1541	19.0 19.1 19.1 19.2 19.2	41.1 41.2 41.3 41.4 41.5	1.1851 1.1856 1.1861 1.1866 1.1872	22.5 22.5 22.6 22.65 22.7	47.6 47.7 47.8 47.9 48.0	1.2196 1.2201 1.2207 1.2212 1.2218	25.9 26.0 26.0 26.1 26.1
28.6 28.7 28.8 28.9 29.0	1.1229 1.1234 1.1239 1.1244 1.1248	15.8 15.8 15.9 15.9	35·3 35·4	1.1546 1.1551 1.1556 1.1561 1.1566	19.3 19.3 19.4 19.4	41.6 41.7 41.8 41.9 42.0	1.1877 1.1882 1.1887 1.1892 1.1898	22.75 22.8 22.9 22.9 23.0	48.1 48.2 48.3 48.4 48.5	1.2223 1.2229 1.2234 1.2240 1.2245	26.2 26.2 26.3 26.35 26.4
29.1 29.2 29.3 29.4 29.5	1.1253 1.1258 1.1263 1.1267 1.1272	16.0 16.1 16.1 16.2 16.25	35.6 35.7 35.8 35.9 36.0	1.1571 1.1576 1.1581 1.1586 1.1591	19.55 19.6 19.65 19.7 19.8	42.1 42.2 42.3 42.4 42.5	1.1903 1.1908 1.1913 1.1919 1.1924	23.0 23.1 23.1 23.2 23.2	48.6 48.7 48.8 48.9 49.0	1.2250 1.2256 1.2261 1.2267 1.2272	26.45 26.5 26.6 26.6 26.7
29.6 29.7 29.8 29.9 30.0	1.1277 1.1282 1.1287 1.1291 1.1296	16.3 16.4 16.4 16.5 16.5	36.1 36.2 36.3 36.4 36.5	1.1596 1.1601 1.1606 1.1611 1.1616	19.8 19.9 19.9 20.0 20.0	42.6 42.7 42.8 42.9 43.0	1.1929 1.1934 1.1940 1.1945 1.1950	23.3 23.4 23.45 23.5	49.1 49.2 49.3 49.4 40.5	1.2278 1.2283 1.2289 1.2294 1.2300	26.7 20.8 26.8 26.9 26.9
30.1 30.2 30.3 30.4 30.5	1.1306	16.6 16.6 16.7 16.7 16.8	36.6 36.7 36.8 36.9 37.0	1.1621 1.1626 1.1631 1.1636 1.1641	20.1 20.1 20.2 20.2 20.3	43.1 43.2 43.3 43.4 43.5	1.1955 1.1961 1.1966 1.1971 1.1976	23.55 23.6 23.7 23.7 23.8	49.6 49.7 49.8 49.9 50.0	1.2305 1.2311 1.2316 1.2322 1.2327	27.0 27.0 27.1 27.1 27.2
30.6 30.7 30.8 30.9 31.0	1.1330	16.8 16.9 17.0 17.0	37.1 37.2 37.3 37.4 37.5	1.1646 1.1651 1.1656 1.1661 1.1666	20.35 20.4 20.5 20.5 20.6	43.6 43.7 43.8 43.9 44.0	1.1982 1.1987 1.1992 1.1998 1.2003	23.8 23.9 23.9 24.0 24.0	50.1 50.2 50.3 50.4 50.5	1.2333 1.2338 1.2344 1.2349 1.2355	27.2 27.3 27.3 27.4 27.4
31.1 31.2 31.3 31.4 31.5	1.1354	17.1 17.2 17.2 17.3 17.3	37.6 37.7 37.8 37.9 38.0	1.1671 1.1676 1.1681 1.1686 1.1692	20.6 20.7 20.7 20.8 20.8	44.1 44.2 44.3 44.4 44.5	1.2008 1.2013 1.2019 1.2024 1.2029	24. I 24. I 24. 2 24. 2 24. 2	50.6 50.7 50.8 50.9 51.0	1.2361 1.2366 1.2372 1.2377 1.2383	27.5 27.55 27.6 27.7 27.7
31.6 31.7 31.8 31.6 32.6	1.1378 1.1383 1.1388	17.4 17.5 17.5	38.2 38.3 38.4	1.1702	20.9 20.9 21.0 21.0 21.1	44.6 44.7 44.8 44.9 45.0	1.2040 1.2045 1.2051	24.35 24.4 24.45 24.5 24.6	51.2		27.8 27.8 27.9 27.9 28.0
32.1 32.2 32.3 32.4 32.5	1.1403	17.7	38.7 38.8 38.9	I.1727 I.1732 I.1737	21.1 21.2 21.3 21.3 21.4		I.2067 I.2072 I.2077	24.6 24.7 24.7 24.8 24.8	51.6 51.7 51.8 51.9 52.0	I.2416 I.2422 I.2427 I.2433 I.2430	28.0 28.1 28.1 28.2 28.2

RELATION OF BRIX, SPECIFIC GRAVITY, AND BAUMÉ—(Continued).

Per Cent of Sugar.	Specific Gravity.	Degree Baumé.	Per Cent of Sugar.	Specific Gravity.	Degree Baumé.	Per Cent of Sugar.	Specific Gravity.	Degree Baumé.	Per Cent of Sugar.	Specific Gravity.	Degree Baumé.
52.1 52.2 52.3 52.4 52.5	1.2444 1.2450 1.2455 1.2461 1.2467	28.3 28.3 28.4 28.4 28.5	58.6 58.7 58.8 58.9 59.0	1.2816 1.2822 1.2978 1.2514 1.2843	31.6 31.7 31.7 31.8 31.8	65.1 65.2 65.3 65.4 65.5	1.3205 1.3211 1.3217 1.3223 1.3229	34.95 35.0 35.05 35.1 35.15	71.6 71.7 71.8 71.9 72.0	1.3610 1.3613 1.3623 1.3629 1.3635	38.2 38.2 38.2 38.3 38.3
52.6 52.7 52.8 52.9 53.0	1.2472 1.2478 1.2483 1.2489 1.2495	28.5 28.6 28.65 28.7 28.75	59.1 59.2 59.3 59.4 59.5	1.2845 1.2851 1.2857 1.2863 1.2869	31.9 31.95 32.0 32.05 32.1	65.6 65.7 65.8 65.9 66.0	1.3235 1.3241 1.3247 1.3253 1.3260	35.2 35.25 35.3 35.35 35.4	72.1 72.2 72.3 72.4 72.5	1.3642 1.3648 1.3655 1.3661 1.3667	38.4 38.4 38.5 38.5 38.6
53.1 53.2 53.3 53.4 53.5	1.2500 1.2506 1.2512 1.2517 1.2523	28.8 28.85 28.9 28.9 29.0	59.6 59.7 59.8 59.9 60.0	1.2875 1.2881 1.2887 1.2893 1.2898	32.15 32.2 32.3 32.3 32.4	66.1 66.2 66.3 66.4 66.5	1.3266 1.3272 1.3278 1.3285 1.3291	35.4 35.5 35.5 35.6 35.6	72.6 72.7 72.8 72.9 73.0	1.3674 1.3680 1.3687 1.3693 1.3699	38.6 38.7 38.7 38.8 38.8
53.6 53.7 53.8 53.9 54.0	1.2529 1.2534 1.2540 1.2546 1.2551	29.1 29.1 29.2 29.2 29.3	60.1 60.2 60.3 60.4 60.5	1.2904 1.2910 1.2916 1.2922 1.2928	32.4 32.5 32.5 32.6 32.6	66.6 66.7 66.8 66.9 67.0	1.3297 1.3303 1.3309 1.3315 1.3322	35.7 35.7 35.8 35.8 35.8	73.1 73.2 73.3 73.4 73.5	1.3705 1.3712 1.3719 1.3725 1.3732	38.9 38.9 39.0 39.0 39.1
54.1 54.2 54.3 54.4 54.5	1.2557 1.2563 1.2568 1.2574 1.2580	29.3 29.4 29.4 29.7 29.5	60.6 60.7 60.8 60.9 61.0	1.2934 1.2940 1.2946 1.2952 1.2958	32.7 32.7 32.8 32.8 32.9	67.1 67.2 67.3 67.4 67.5	1.3327 1.3334 1.3340 1.3346 1.3352	35.9 36.0 36.0 36.1 36.1	73.6 73.7 73.8 73.9 74.0	1.3738 1.3745 1.3751 1.3757 1.3764	39.1 39.2 39.2 39.3 39.3
54.6 54.7 54.8 54.9 55.0	1.2585 1.2591 1.2597 1.2602 1.2608	29.6 29.6 29.7 29.7 29.8	61.1 61.2 61.3 61.4 61.5	1.2964 1.2970 1.2975 1.2981 1.2987	32.9 33.0 33.0 33.1 33.1	67.6 67.7 67.8 67.9 68.0	1.3359 1.3365 1.3371 1.3377 1.3384	36.2 36.3 36.3 36.4	74.1 74.2 74.3 74.4 74.5	1.3770 1.3777 1.3783 1.3790 1.3796	39 · 4 39 · 4 39 · 5 39 · 5 39 · 6
55.1 55.2 55.3 55.4 55.5	1.2614 1.2620 1.2625 1.2631 1.2637	29.8 29.9 29.9 30.0 30.05	61.6 61.7 61.8 61.9 62.0	1.2993 1.2999 1.3005 1.3011 1.3017	33.2 33.3 33.3 33.4	68.1 68.2 68.3 68.4 68.5	1.3390 1.3396 1.3402 1.3408 1.3415	36.4 36.5 36.5 36.6 36.6	74.6 74.7 74.8 74.9 75.0	1.3803 1.3809 1.3816 1.3822 1.3828	39.6 39.7 39.7 39.8 39.8
55.6 55.7 55.8 55.9 56.0	1.2642 1.2648 1.2654 1.2660 1.2665	30.1 30.15 30.2 30.25 30.3	62.1 62.2 62.3 62.4 62.5	1.3023 1.3029 1.3035 1.3040 1.3047	33.4 33.5 33.5 33.6 33.6	68.6 68.7 68.8 68.9 69.0	1.3421 1.3427 1.3433 1.3440 1.3446	36.7 36.7 36.3 36.8 36.9	75.1 75.2 75.3 75.4 75.5	1.3835 1.3842 1.3848 1.3855 1.3861	39.9 39.9 40.0 40.0
56.1 56.2 56.3 56.4 56.5	1.2671 1.2677 1.2683 1.2688 1.2694	30.4 30.4 30.5 30.5 30.6	62.6 62.7 62.8 62.9 63.0	1.3053 1.3059 1.3065 1.3071 1.3077		69.1 69.2 69.3 69.4 69.5	1.3452 1.3458 1.3405 1.3471 1.3477	36.9 37.0 37.0 37.1 37.1	75.6 75.7 75.8 75.9 76.0	1.3868 1.3874 1.3880 1.3887 1.3894	40.1 40.2 40.2 40.3 40.3
56.6 56.7 56.8 56.9 57.0	1.2700 1.2706 1.2712 1.2717 1.2723	30.6 30.7 30.7 30.8 30.8	63.1 63.2 63.3 63.4 63.5	1.3083 1.3089 1.3095 1.3101 1.3107	33.9 34.0 34.0 34.1 34.1	69.6 69.7 69.8 69.9 70.0	1.3484 1.3490 1.3496 1.3502 1.3509	37 · 2 37 · 3 37 · 3 37 · 3	76.1 76.2 76.3 76.4 76.5	1.3900 1.3907 1.3913 1.3920 1.3926	40.4 40.4 40.5 40.5 40.6
57.1 57.2 57.3 57.4 57.5	1.2740	30.9 30.9 31.0 31.0	63.6 63.7 63.8 63.9 64.0	1.3113 1.3119 1.3126 1.3132 1.3138		70.1 70.2 70.3 70.4 70.5	1.3515 1.3521 1.3528 1.3534 1.3540	37.4 37.5 37.5 37.6 37.6	76.6 76.7 76.8 76.9 77.0	1.3933 1.3940 1.3946 1.3953 1.3959	40.6 40.7 40.7 40.8 40.8
57.6 57.7 57.8 57.9 58.0	1.2764	31.1 31.2 31.2 31.3	64.1 64.2 64.3 64.4 64.5	1.3144 1.3150 1.3156 1.3162 1.3168	34.4 34.5 34.5 34.6 34.6	70.6 70.7 70.8 70.9 71.0	1.3546 1.3553 1.3559 1.3565 1.3572	37 · 7 37 · 7 37 · 8 37 · 8 37 · 8	77.1 77.2 77.3 77.4 77.5	1.3966 1.3972 1.3979 1.3986 1.3992	40.8 40.9 41.0 41.0 41.0
58.1 58.2 58.3 58.4 58.5	1.2793 1.2799 1.2804	31.4 31.5 31.5 31.5	64.6 64.7 64.8 64.9 65.0	1.3174 1.3180 1.3186 1.3192 1.3198	34.7 34.7 34.8 34.8 34.9	71.1 71.2 71.3 71.4 71.5	1.3578 1.3585 1.3591 1.3597 1.3604	38.0 35.1	77.6 77.7 77.8 77.9 78.0	1.3999 1.4005 1.4012 1.4019 1.4025	41.1 41.1 41.2 41.2 41.3

				,							
Per Cent of Sugar.	Specific Gravity.	Degree Baumé.	Per Cent of Sugar.	Specific Gravity.	Degree Baumé.	Per Cent of Sugar.	Specific Gravity.	Degree Baumé.	Per Cent of Sugar.	Specific Gravity.	Degree Baumé.
78.1	1.4032	41.3	80.1	1.4165	42.3	82.1	1.4300	43·3	84.1	I.4437	44.2
78.2	1.4039	41.4	80.2	1.4172	42.3	82.2	1.4307	43·3	84.2	I.4443	44.3
78.3	1.4045	41.4	80.3	1.4179	42.4	82.3	1.4314	43·4	84.3	I.4450	44.3
78.4	1.4052	41.5	80.4	1.4185	42.4	82.4	1.4320	43·4	84.4	I.4457	44.3
78.5	1.4058	41.5	80.5	1.4192	42.5	82.5	1.4327	43·5	84.5	I.4464	44.4
78.6	1.4065	41.6	80.6	1.4199	42.5	82.6	1.4334	43.5	84.6	1.4471	44.4
78.7	1.4072	41.6	80.7	1.4205	42.6	82.7	1.4341	43.5	84.7	1.4478	44.5
78.8	1.4078	41.7	80.8	1.4212	42.6	82.8	1.4348	43.6	84.8	1.4485	44.5
78.9	1.4085	41.7	80.0	1.4219	42.7	82.9	1.4354	43.6	84.9	1.4492	44.6
79.0	1.4092	41.8	81.0	1.4226	42.7	83.0	1.4351	43.7	85.0	1.4498	44.6
79.1	1.4098	41.8	81.1	1.4232	42.8	83.1	1.4368	43.7	85.1	1.4505	44.7
79.2	1.4105	41.9	81.2	1.4239	42.8	83.2	1.4375	43.8	85.2	1.4512	44.7
79.3	1.4112	41.9	81.3	1.4246	42.0	83.3	1.4382	43.8	85.3	1.4519	44.8
79.4	1.4119	42.0	81.4	1.4253	42.0	83.4	1.4388	43.9	85.4	1.4526	44.8
79.5	1.4125	42.0	81.5	1.4259	43.0	83.5	1.4395	43.9	85.5	1.4533	44.9
79.6	1.4132	42.I	81.6	1.4266	43.0	83.6	1.4402	44.0	85.6	1.4540	44.9
79.7	1.4138	42.I	81.7	1.4273	43.1	83.7	1.4409	44.0	85.7	1.4547	45.0
79.8	1.4145	42.2	81.8	1.4280	43.1	83.8	1.4410	44.1	85.8	1.4554	45.0
79.9	1.4152	42.2	81.9	1.4287	43.2	83.9	1.4423	44.1	85.9	1.4561	45.1
80.0	1.4158	42.2	82.0	1.4293	43.2	84.0	1.4430	44.2	86.0	1.4568	45.1

RELATION OF BRIX, SPECIFIC GRAVITY, AND BAUMÉ-(Concluded).

extreme accuracy is required, the double dilution method of Wiley should be employed, which makes due allowance for the volume of the precipitate.

Double Dilution Method.*—Take half the normal weight of the sample and make up the solution to 100 cc., using the appropriate clarifier. Take the normal weight of the sample and make up a second solution with the clarifier to 100 cc. Filter and obtain direct polariscopic readings of both solutions. Invert each in the usual manner and obtain the invert reading of the two.

The true direct polarization of the sample is the product of the two direct readings divided by their difference. The true invert polarization is the product of the two invert readings divided by their difference.

Determination of Raffinose in Beet Sugar Molasses.—For the determination of sucrose and raffinose when present in the same solution, use the following formulas of Creydt: †

Sucrose =
$$\frac{0.5188a - b}{0.8454}$$
 (1)

Raffinose =
$$\frac{a-S}{1.85}$$
, (2)

where a=direct reading, b=reading after inversion, and S=per cent of sucrose.

^{*} Wiley and Elwell, Analyst, 1896, 21, p. 184.

[†] U. S. Dept. Agric., Bur. of Chem., Bul. 107 (rev.), p. 43.

Davoll * recommends for purposes of clarification of the molasses the use of powdered zinc after inversion of the molasses sample according to Clerget's method. He adds I gram of the zinc to the sample after inversion while at the temperature of 69° C., allowing it to act for three to four minutes at that temperature, after which he cools and filters, with the production of an almost colorless solution.

Determination of Reducing Sugar.—(Estimated as Dextrose.)—Dilute 5 grams of molasses or syrup with water in a 100-cc. graduated flask, using 2 to 5 cc. of subacetate of lead solution. Make up to 100 cc., filter, take an aliquot part of the filtrate (25 to 50 cc.) and make this up to 100 cc., the amount taken being such that, when diluted, the solution will contain not more than \frac{1}{2}\%0 of dextrose. If lead subacetate has been used to clarify, add to the aliquot part taken and before dilution, enough sodium sulphate to precipitate the excess of lead, then filter and make up to the 100 cc. mark.

Determine the reducing sugar in this solution by either volumetric or gravimetric Fehling processes.

U. S. Standard Molasses is molasses containing not more than 25% of water, nor more than 5% of ash.

Adulteration of Molasses and Syrups.—A common adulterant of all these products is commercial glucose. From its water-white color and inert sweetness, no less than from its cheapness, it forms an admirable adulterant for dark-colored or low-grade molasses and syrups, counteracting to a great extent by its smoothness the strong and often disagreeable taste of the inferior products with which it is mixed. Thus a grade of molasses too cheap to be ordinarily used for food purposes can be made to assume the appearance, and to some extent the taste, of the higher-priced and light-colored grades, by admixture with commercial glucose.

Tin salts are also used to improve the color of low-grade or dark molasses, and bleaching agents, such as sulphurous acid, are frequently employed. Copper is sometimes found, due to utensils or vessels used in processes of manufacture.

Lead may occur in maple syrup, due to the leaden plugs or spigots through which the sap is sometimes drawn from the trees.

Detection and Determination of Commercial Glucose.†—From the direct polarization of a normal solution of molasses or syrup the presence

^{*} Jour. Am. Chem. Soc., 25 (1903), p. 1019.

[†] Leach, ibid., p. 982.

or absence of commercial glucose can usually be established. The direct polarization of a normal solution of pure molasses should not be much in excess of 50° on the Soleil-Ventzke scale, while a pure, dark-colored molasses should polarize well under 40°. Golden syrup and maple syrup read higher than molasses, and a normal solution of pure maple syrup may have a direct polarization as high as 65°, being more often than not above 60°.

An excessively high direct polarization is at once an indication of the presence of commercial glucose, while an invert reading at ordinary room temperature to the right of the zero-point is an almost positive proof of its presence in either of the above products.

The optically active constituents of commercial glucose, viz., dextrin, maltose, and dextrose, are present in such varying amounts, that it is impossible to determine accurately the exact amount of this adulterant in complex saccharine products which themselves contain components common to glucose. Its approximate amount can, however, be very satisfactorily estimated in molasses and syrups by the use of the following formula:

$$G = \frac{(a-S)100}{175}, *$$

where G-per cent of commercial glucose, a-direct polarization, and S-per cent of cane sugar previously obtained from the Clerget formula (p. 588). A large amount of invert sugar present affects the accuracy of this formula. It is especially applicable to maple syrup, wherein the per cent of invert sugar is small, but may be applied also to molasses and golden syrup, wherein the amount of invert sugar is not so large but that results may be obtained as close as could be expected from an empirical formula.†

In saccharine products containing considerable invert sugar the invert reading at 87° C. obtained as directed on page 639, is divided by

^{*} Leach, U. S. Dept. of Agric., Bur. of Chem., Bul. 65, p. 48.

[†] This formula is based on the assumption that 42° Bé. mixing glucose, the grade specially made and used for admixture with molasses, syrups, and honey, has a maximum polarization of 175° V. It was adopted as a result of investigations made some years ago by the author, but subsequently it appeared that 42° Bé. mixing glucose polarizes lower than formerly. Thus a sample recently examined by the author polarized at 162.4° V. Pending further investigations it seems best for the present to retain the old formula, for, while it undoubtedly gives low results, especially with higher admixtures of glucose, it approximates the truth more closely than would be expected, perhaps because it tends to compensate for the error due to substances in genuine molasses and honey that polarize to the right after inversion. Furthermore, it has been adopted by the A. O. A. C. To avoid misunderstanding, express results in terms of glucose polarizing at that factor.

the appropriate factor (163) to obtain the percentage of commercial glucose.

While theoretically pure molasses and syrups would be expected to show no rotation when polarized at 87° C. after inversion, as a matter of fact most samples exhibit a decidedly right-handed reading at that temperature. Occasionally a zero reading is noted, and in rare instances a slight left-handed rotation occurs under the above conditions.

Dextro-rotation is undoubtedly caused by some form of decomposition or fermentation. It may be due to a preponderance of dextrose in the reducing sugars, since levulose is more easily decomposed than dextrose, or it may be caused by the decomposition products formed when the raw juice is being defecated with lime, or again it might result from a special fermentation forming dextran.

The following table shows results by A. H. Bryan* of polarization of samples of Louisiana molasses and syrup of known purity, showing especially the invert readings at 87° C.:

POLARIZATION	OF	LOUISIANA	MOI ASSES	AND	CVDIID
PULAKIZATION	Ur	LOUISIANA	MOLASSES	AND	SYRUP.

	MOI	ASSES.			SY	RUP.	
Direct Polariza- tion		d Invert	Dry Substance.	Direct Polariza- tion	Correcte Polariz	d Invert	Dry Substance
at 20° C.	At 20° C.	At 87° C.	oubstance.	at 20° C.	At 20° C.	At 87° C.	Substance
• V.	• v.	° V.	Per Cent.	° V.	° V.	° V.	Per Cent.
40.8	- 20.24	+2.2	80.8	48.4	-17.6	+1.98	74-3
24.6	-20.9	+2.2	76.8	54.0	- 18.7	+3.30	68.3
26.0	- 18.26	+3.52	76.8	50.2	-12.1	+6.16†	ŀ
42.4	- 16.94	+2.42	78.2	50.4	-14.3	+1.76	
52.4	-16.28	+2.20	69.1	61.8	-16.5	+2.20	1
55.6	- 13.59	+4.18	69.6	(l
39.6	- 18.04	+2.20	80.8				
39.6	-17.82	+2.20	79.0	Average.		+2.65	
44.0	-17.16	+2.64	72.0	Maximu	m	+6.16	
42.0	-17.60	+2.42	73.8 76.1	Minimur	n	0.00	
42.4 41.6	- 17.27 - 16.94	+3.52 +3.96		1		-	-
52.4	-10.94 -17.60	+3.52	74.0 76.1	1			
26.6	-19.8	0.00	78.1				
50.8	- 25.c8	+1.10	87.5	l			
22.6	- 16.72	+3.96	84.1	[
41.6	-14.74	+1.10	75.0				
45.6	-15.4	+2.20	78.0	ļ			

^{*} A. O. A. C. Proc., 1908, U. S. Dept. of Agric., Bur. of Chem., Bul. 122, p. 182.

[†] Sample ropy and badly fermented.

TYPICAL	ANALYSES	OF	MOLASSES	AND	SYRUPS	ADULTERATED	WITH
			COMMERC	IAL G	LUCOSE.		

	P	olarizatio	n.		ي			
	Direct.	Invert.	Tempera- ture.	Per Cent Sucrose (Clerget's Formula).	Reducing Sugar (Der trose).	Commercial Glucose (Leach's Formula).	Moisture.	Ash.
(a) Molasses. (b) " (c) " (a) Golden drip syrup. (b) " (c) " (a) Maple syrup. (b) " (c) " (a) Machine syrup. (c) " (d) Machine syrup. (d) " (e) " (f)	62 98-7 109-7 73-5 109-4 143-6 76-3 77-9 87-0	+36.3 +71.9 +90 +39.8 +87.6 +136.0 +7.6 +24 +30.6	18° 17° 18° 17° 18.4° 18.6°	19.9 14.5 25 16.9 5.6 51 40.1	30.03 27.62 33.11 31.61 33.44 38.17 10.55	24.6 45.0 54.4 27.7 52.8 78.5 14.4 21.6	29.36 27.98 22.02 23.67 24.48 21.52 31.91 23.44 28.80	3.83 3-53 2.67 3-94 2.51 1.00 0.65

Determination of Dextrin. — According to Beckman's method a weighed amount of the honey or molasses is diluted with an equal volume of water and from ten to twelve times its volume of methyl alcohol is added. The precipitated dextrin is collected in a tared filter and thoroughly washed with methyl alcohol, after which it is dried and weighed.

Reduction of Saccharine Products to an Ash for Mineral Analysis.—If a considerable quantity of molasses, syrup, or other saccharine substance is to be burnt to an ash, it is both tedious and annoying to ignite directly, by reason of the excessive swelling and frothing of such substances during ignition. Small quantities of molasses, syrup, or honey may with care be reduced to an ash by the method described on page 586.

If a readily controlled electric current is available, it may be utilized as follows:* Mix 100 grams of molasses, syrup, or other saccharine solution, which should be evaporated to syrupy consistency if not already such, with about 35 grams of concentrated sulphuric acid in a large porcelain evaporating-dish. An electric current is then passed through it while stirring, by placing one platinum electrode in the bottom of the dish near one side and attaching the other to the lower end of the glass rod, with which the contents are stirred. Begin with a current of about 1 ampere and gradually increase to 4.† In from ten to fifteen minutes

^{*} Leach, 32d An. Rept. Mass. State Board of Health (1900), p. 653. Reprint, p. 37. This method is preferred to the ordinary method of heating with sulphuric acid, especially in case of molasses, because, if properly manipulated, it so quietly comes into the form of a very finely divided char or powder, especially adapted for subsequent quick ignition.

[†] Modified from method of Budde and Schou for determining nitrogen electrolytically. Ztschr. anal. Chem., 38 (1899), p. 345.

the mass is reduced to a fine, dry char, which may then be readily burnt to a white ash in the original dish over a free flame or in a muffle.

Or, 100 grams of the molasses or syrupy solution to be ashed may be first evaporated to dryness and afterward mixed with from 10 to 20 cc. of concentrated sulphuric acid in a porcelain evaporating-dish, or if the substance to be ashed be a dry sugar or confectionery, 20 grams are mixed with the above amount of acid. Heat is gently applied by means of the gas flame till the swelling and frothing have ceased, which usually requires only a few minutes. The final ignition is then accomplished in the usual manner, nitric acid being added if necessary to completely destroy the organic matter.

Determination of Tin in Molasses.—Fuse the ash from a weighed portion of the sample with sodium hydroxide in a silver crucible, dissolve in water, and acidulate with hydrochloric acid; filter and precipitate the tin from this solution with hydrogen sulphide; wash the precipitate on a filter and dissolve it in an excess of ammonium sulphide. Filter this solution into a tared platinum dish, and deposit the tin directly in the dish by electrolysis, using a current of 0.05 ampere and the apparatus described on page 608.

Distinction between Invert Sugar, Maltose, and Lactose.*—All these sugars reduce Fehling's solution. Dextrose and levulose (invert sugar) when boiled with Barfoed's copper acetate solution (14 grams crystallized copper acetate and 5 cc. acetic acid in 200 cc. water) will form a precipitate of cuprous oxide, while neither maltose nor lactose will do this. The solution, which has thus been tested for invert sugar and found to be free, or the filtrate from the cuprous oxide precipitate, is treated with an excess of basic lead acetate, filtered, and to the filtrate is added an excess of sodium sulphate solution to precipitate the lead. The solution is again filtered and treated with copper sulphate solution, if not already blue. It is then made alkaline with sodium hydroxide and heated to boiling. A red precipitate of cuprous oxide at this stage indicates either lactose or maltose or both.

A solution of the sugar, made strongly ammoniacal, is then mixed with alkaline bismuth solution † and the container is set in a waterbath at 60° C. Maltose soon reduces the bismuth, but lactose does not.

To test for lactose, add strong nitric acid to the solid sugar residue

^{*} Bartley and Mayer, Merck's Report, 12 (1903), p. 100.

[†] This reagent is prepared as follows: Bismuth subnitrate, 2 grams; Rochelle salt, 4 grams; sodium hydroxide, 8 grams; dissolved in 100 cc. of water by the aid of heat.

and warm gently till red fumes come off. Then set the container in hot water and cool gradually. Crystals of mucic acid appear after a time if any appreciable amount of lactose be present.

Determination of Lactose or Maltose.—Either sugar, if in solution free from other reducing sugars, may be determined by the volumetric Fehling method (p. 591) or by the Defren method, using the table on page 595.

For the determination of maltose in commercial glucose, see page 630. Estimation of Cane Sugar and Dextrose in Mixtures.—Obtain true direct and invert readings of a normal solution of the mixture. Determine the per cent of sucrose by Clerget's formula (p. 588). This figure represents the right-handed rotation due to sucrose. Subtracting this from the direct polarization, the difference represents the right-handed rotation due to dextrose. The specific rotary power of sucrose is 66.5 and that of dextrose 52.3.

Calling d the percentage of dextrose and R' the right-handed rotation due to dextrose as above obtained, if the Soleil-Ventzke scale is used,

$$66.5:52.3=d:R'$$
,

whence

$$d = \frac{66.5R'}{5^2 \cdot 3}.$$

Determination of Levulose.*—On page 589 attention was called to the variation in the rotary power of levulose with the temperature. This variation is constant, and I gram of levulose in 100 cc. of water produces a decrease in left-handed reading of 0.0357° on the cane sugar (Ventzke) scale for each I°C. increase in temperature. Therefore, the weight of levulose present in a given solution can be calculated from the polariscopic readings at two temperatures, using a water-jacketed tube, as described on page 639.

$$L = \frac{R - R'}{0.0357 (t - t')},$$

where

L=weight of levulose,

R=reading at higher temperature t,

R'=reading at lower temperature t.

The percentage of levulose present in the solution may readily be calculated as follows:

^{*} Wiley, Agric. Anal., p. 272.

If

L' = percentage of levulose,

L = weight of levulose in solution,

W = weight of sugar sample made up to 100 cc.,

$$L' = \frac{L \times 100}{W}.$$

In a normal solution W = 26.048.

ANALYSIS OF MAPLE PRODUCTS.

Determination of Moisture.—This is accomplished by direct drying with sand, or by calculation from the specific gravity, or, preferably from the refractive index. See molasses methods, page 613.

Determination of Ash.—Burn 5 grams in a platinum dish by the usual method, observing the precautions given for molasses, page 614.

Soluble and Insoluble Ash.*—To the platinum dish containing the ash add 40 cc. of hot water and boil gently for two minutes. Filter through a small ashless filter, and wash with hot water until the filtrate amounts to 100 cc. Return the filter to the dish used for ashing, burn at a low red heat, cool and weigh, thus obtaining the insoluble ash. The soluble ash is obtained by difference, subtracting the weight of insoluble from that of total ash.

Alkalinity of Soluble Ash.†—Allow the filtrate from the above determination to cool, then titrate with tenth-normal hydrochloric acid, using methyl orange as an indicator.

Alkalinity of Insoluble Ash.†—Add excess of tenth-normal hydrochloric acid (usually 10 to 15 cc.) to the ignited insoluble ash in the platinum dish, heat to the point of boiling over an asbestos plate, allow to cool, and titrate excess of hydrochloric acid with tenth-normal sodium hydroxide, using methyl orange as an indicator.

Express the alkalinity in each case as the number of cubic centimeters of tenth-normal acid used on the ash of 1 gram of sample.

Polarization.—See page 614.

Determination of Reducing Sugar.—See page 621.

Determination of Malic Acid Value.—Modified Leach and Lythgoe Method.‡—Weigh 6.7 grams of the sample into a 200 cc. beaker, and add

^{*} A. H. Bryan, U. S. Dept. of Agric., Bur. of Chem., Circ. No. 40, p. 6.

[†] U. S. Dept. of Agric, Bur. of Chem., Bul. 107 (rev.), p. 69.

[‡] Jour. Am. Chem. Soc., 26, 1904, pp. 380 and 1536; U. S. Dept. of Agric., Bur. of Chem., Bul. 107 (rev.), p. 74.

water to make a volume of 20 cc. Add 2 drops of ammonium hydroxide (specific gravity, 0.90), 1 cc. of a 10% solution of calcium chloride, and 60 cc. of 95% alcohol. Cover the beaker with a watch glass, heat for one-half hour on a water bath, then turn off the flame and allow the beaker to stand overnight. Filter the material in the beaker through good quality filter paper, wash the precipitate with hot 75% alcohol until the filtrate measures 100 cc., dry and ignite. Add from 15 to 20 cc. of tenth-normal hydrochloric acid to the ignited residue, thoroughly dissolve the lime by heating carefully to just below boiling, cool and titrate the excess of acid with tenth-normal sodium hydroxide, using methyl orange as an indicator. One-tenth of the number of cubic centimeters of acid neutralized by the ignited residue expresses the malic acid value. Run blank determinations on reagents, using the same amounts, particularly of ammonium hydroxide, as were used in the original determination, and make the necessary correction.

Determination of Lead Number. — Winton Method.* — Weigh 25 grams of the material (or 26.048 grams if it is desired to determine sugars polariscopically in the same portion) and transfer by means of water into a 100-cc. flask. Add 25 cc. of standard lead subacetate solution, fill to the mark, shake, allow to stand at least three hours and filter through a dry filter. From the clear filtrate pipette off 10 cc., dilute to 50 cc., add a moderate excess of sulphuric acid, and 100 cc. of 95% alcohol. Let stand over night, filter on a Gooch crucible, wash with 95% alcohol, dry at a moderate heat, ignite at low redness for three minutes, taking care to avoid the reducing cone of the flame, cool, and weigh. Calculate the amount of lead in the precipitate, using the factor 0.6829, subtract this from the amount of lead in 2.5 cc. of the standard solution, multiply the remainder by 100, and divide by 2.5, thus obtaining the lead number.

The standard lead subacetate is prepared by diluting one part of the ordinary solution (page 586) with four volumes of water, filtering if not clear. It is standardized by determining the lead in 25 cc. as above described. The solution deposits a slight precipitate on standing, but this does not usually appreciably affect its strength.

Determination of Hortvet Number.†—The method depends on the principle that the volume of the precipitate, by treatment of the sugar solution or syrup under fixed conditions with alumina cream and subacetate of lead, varies with the amount of refined sugar present.

^{*} Jour. Am. Chem. Soc., 28, 1906, p. 1204.

[†] Ibid., 26, 1904, p. 1532.

Apparatus.—(1) A tube, adapted to be carried in the shield of the centrifuge. This tube, which is 15.3 cm. in length, has a wide cylindrical portion 3 cm. in diameter, narrowed at the top to a neck 2 cm. in diameter, and at the bottom to a stem graduated in tenths to 5 cc.

(2) A holder, made of pine or white wood, of a size adapted to carry the tube in the shield of the centrifuge. The holders and tubes should be arranged in balanced pairs in the centrifuge.

Procedure.—Introduce 5 cc. of syrup or 5 grams of sugar into the tube. Add 10 cc. of water, and dissolve completely. Next add 10 drops of alumina cream, and 1.5 cc. of lead subacetate. Shake thoroughly, and allow to stand from forty-five to sixty minutes. Place the tube in its holder in the centrifuge shield, and run six minutes. If, after the end of this time, any material adheres to the sides of the wide part of the tube, loosen with a small wire or by giving the tube a slight twist, then run the tube six additional minutes, and finally read the volume of the precipitate in the stem, estimating to 0.01 cc.

Run a blank with the above reagents in water, subtracting the blank reading from that of the precipitate. In the case of syrup, reduce to the 5-gram basis by dividing by the specific gravity of the sample. If the sugar content of the sample is known, the specific gravity can be calculated from the table on page 617. For pure maple syrup 1.33 is very nearly correct.

The following table shows results obtained by Hortvet on two samples of maple syrup of known purity, mixed with varying amounts of refined cane sugar syrup of the same density:

Purity.	Five Minutes.	Com- puted Precipi- tate.	Differ- ence.	Ten Minutes.	Com- puted Precipi- tate.	Differ- ence.	Twelve Minutes.	Com- puted Precipi- tate.	Differ- ence.
Per Cent.	cc. 4.90	ec.	cc.	cc. 3.29	cc.	ec.	ec. 2.80	œ.	cc.
25	1.00	I.22	0.22	0.70	0.82	0.12	0.65	0.70	0.05
50	3.60	2.45	1.15	1.50	1.54	0.14	1.40	1.40	0.00
75	4.60	3.67	0.93	2.77	2.47	0.30	1.90	2.10	0.20
100	5.40			4.40			3.75		
10	0.30	0.54	0.24	0.28	0.44	0.16	0.28	0.37	0.00
20	1.27	1.08	0.19	0.80	0.88	0.70	0.70	0.75	0.05
60	5.00	3.24	1.76	2.70	2.64	0.06	1.70	2.25	0.55

The blank at the end of twelve minutes was 0.44 cc. The machine used for the above experiment had a radius of 18.5 cm. and a speed of 1600 revolutions per minute. Results obtained by Hortvet on known

pure maple syrups vary from 1.2 cc. to about 2.5 cc., and on known pure maple sugars from 1.8 cc. to 4 cc.

Commercial brands of adulterated syrups and sugars give such precipitates as 0.00 cc., 0.02 cc., 0.05 cc., and 0.08 cc. Hortvet regards with suspicion a syrup testing lower than 1.2 cc., and when the result is below 1 cc., the sample is positively condemned as being mixed with refined cane sugar. In the case of sugar, a somewhat higher minimum figure is adopted than with syrup. In view of the fact that the speed has much to do with the volume of the precipitate, the analyst should make a series of similar experiments with his own centrifuge, and work out his own standards. Results may be better compared with each other, if calculated on the water-free basis.

In case of doubt, and in fact in all cases at first, it would be well to make confirmatory tests, such as determining the ash and reducing sugar.

Sy's Lead Method.*—In a 25-cc. graduated cylinder introduce 5 cc. of syrup, or 5 grams of sugar which is afterwards dissolved in a little water. Add water to the 15 cc. mark and 2 cc. of lead subacetate solution. Shake thoroughly and allow the mixture to stand twenty hours. Then read the volume of the precipitate, which for pure maple products should be at least 3 cc. and is usually over 5 cc.

ANALYSIS OF COMMERCIAL GLUCOSE.

Wiley† has worked out a method for calculating the percentage of dextrin, maltose, and dextrose present in commercial glucose, based on the specific rotary power of these substances and on the reducing power of maltose and dextrose. To apply this method, the operator, if he has a polariscope reading in sugar scale degrees, must ascertain the equivalent readings in angular degrees from the table on page 583, and calculate the specific rotary power in each case from the formula

$$(\alpha)_D = \frac{100a}{cl}$$
, page 584.

Thus, if he possesses a Schmidt and Haensch instrument, he should multiply the true reading, as obtained on that instrument, with a normal

^{*} Jour. Am. Chem. Soc., 30, 1908, p. 1430.

[†] Chem. News, 46, p. 175; Agric. Anal., 3, pp. 288-290.

solution of the given sugar or mixture, by the factor 0.3468, to convert the reading into circular degrees from which to figure the specific rotary power as above.

The specific rotary power of dextrin is fixed at 193, that of maltose 138, and that of dextrose at 53.

Then if P=total polarization of the mixture in terms of specific rotary power, d=per cent dextrose, m=per cent maltose, and d'=per cent dextrin,

$$P = 53d + 138m + 193d'$$
. (1)

The value of P is obtained from observation and calculation as above described on a known solution of the sample, say 10 grams in 100 cc. The reducing sugars, maltose and dextrose, are then removed, preferably by oxidation with cyanide of mercury, as follows:*

Prepare the reagent by dissolving 120 grams mercuric cyanide and 120 grams sodium hydroxide in water, mixing the two solutions, and making up to 1,000 cc. Remove any precipitate that may gather by filtration.

Make a solution of 10 grams of the glucose sample in 100 cc. and take 10 cc. of this solution in a 50-cc. graduated flask. Add sufficient mercuric cyanide solution to have an excess of reagent after the oxidation (from 20 to 25 cc.), and boil for three minutes under a hood with a good draft. Cool and neutralize the alkali with concentrated hydrochloric acid, adding the latter till the brown color is discharged. By this method the optical activity of the maltose and dextrose is discharged, while that of the dextrin remains unaffected. From the polariscope reading calculate as above the specific rotary power of the dextrin (P'). Then

$$P' = 193d'$$
. (2)

The reducing power on Fehling's solution of dextrose is to that of maltose as 100 is to 62. Whence, if R=reducing sugar (reckoned as dextrose) we have

$$R = d + 0.62m$$
. (3)

Subtracting equation (2) from equation (1) we have

$$P-P'=53d+138m$$
. (4)

^{*} Wiley, Agric. Analysis, p. 290.

Multiplying equation (3) by 53 and subtracting from equation (4),

$$P-P'=53d+138m$$
,
 $53 R=53d+32.86m$,
 $P-P'-53R=105.14m$ (5)

Therefore

$$d=R-0.62m, \quad \ldots \quad \ldots \quad (7)$$

$$d' = \frac{P'}{193}. \qquad (8)$$

Determination of Dextrin in Commercial Glucose.—One volume of the sample is well shaken with about 10 volumes of 90% alcohol, and the precipitated dextrin is separated by filtration through a tared filter, washed thoroughly with strong alcohol, dried at 100°, and weighed.

Qualitative Tests for Commercial Glucose. — Several confirmatory chemical tests may be employed for commercial glucose, aside from the optical test with the polariscope. Thus a precipitate of dextrin by treatment of the sample with an excess of strong alcohol, in the absence of mineral salts insoluble in alcohol, is strongly indicative of commercial glucose. An excess of calcium sulphate in the ash also points strongly to the presence of glucose.

Arsenic in Commercial Glucose.—Like all products wherein commercial sulphuric acid is employed in its manufacture, glucose sometimes contains arsenic, though usually in minute traces. Arsenic is readily indicated, when present, by the Gutzeit test, conducted as follows: 2 grams of the sample are introduced into a small Erlenmeyer flask of about 100 cc. capacity, and diluted with 5 to 10 cc. of water. Scraps of arsenicfree, granulated zinc are then added. A small filter-paper is carefully folded smoothly around the bottom of a cork that loosely fits the mouth of the flask, and is moistened with a concentrated solution of mercuric chloride. From 6 to 8 cc. of arsenic-free concentrated hydrochloric* or sulphuric acid are then added to the flask, so as to produce rapid, but not too violent evolution of gas, and the cork is loosely inserted.

After ten minutes the cork is removed, and, if a yellow stain is present on the filter, arsenic is indicated. The amount of arsenic present varies

^{*} Hydrochloric acid is better than sulphuric acid, as the action is much more brisk with pure zinc.

with the depth of color, and if a large amount is present the stain may be dark brown or even black.

Sulphides interfere with the Gutzeit test, but are rarely present in commercial glucose. Unless sure of the purity of the reagents it is well to make a blank test thereon. In such a blank, the filter should be perfectly white after ten minutes.

The amount of arsenic may be roughly determined colorimetrically by the Gutzeit method.

For more careful determination, employ the Marsh apparatus, into which the diluted glucose may be directly introduced without previous treatment.

HONEY.

Composition and Occurrence.—Honey is the saccharine product deposited by bees (Apis mellifica and A. dorsata) in the cells of honey comb, which the insect forms out of wax secreted by its body. Honey has its source chiefly in the nectaries of flowers, from which the bees abstract it, also in the juices of ripe fruits and the exudations of leaves (honeydew). While in the honey-sac of the bee, the sucrose, which forms the chief constituent of the fruit juice or nectar, becomes for the most part inverted, forming, in the honey, dextrose and levulose. The evaporation to a syrupy consistency is effected in the hive by exposure to a current of air, produced by fanning of the wings of the bees.

The flavor of honey varies considerably, according to its source. Besides water, the sugars, and mineral matters, pollen is usually present, derived from the flowers, also as a rule a small quantity of wax, and nearly always appreciable amounts of various organic acids, such as formic.

European Honey.—Neufeld * gives the following limits for pure honey:

Water.	8.30 to 33.59%
Protein	0.03 " 2.67%
Invert sugar	49.59 " 93.96%
Sucrose	0.10 " 10.12%
Dextrin	0.99 " 9.70%
Formic acid	0.03 " 0.21%
Ash	0.02 " 0.68%

^{*} Der Nahrungsmittelchemiker als Sachverständiger., Berlin, 1907, p. 275.

Canadian Honey.—A large number of samples of genuine honey analyzed in 1897 for the Department of Inland Revenue, Canada (Bul. 47), showed the following variations:

Direct polarization	- 2.4	to	-19
Invert "	-10.2	"	-28
Sucrose (by Clerget)	0.5	"	7.64%
Invert sugar	60.37	"	78.8%
Water	12 .	"	33%
Ash	0.03	"	0.50%

American Honey.—Browne* has examined 97 samples of American and Hawaiian honey, representing the product made from the nectar of numerous flowers as well as honeydew. Maxima and minima of polarizations and analyses of some of the more important kinds, and of all the levorotatory and the dextrorotatory samples are given in the table on page 635.

As regards the chemical characteristics of honey from different flowers, Browne states that alfalfa honey usually has less dextrin and undetermined matter—the so-called 'impurities"—and more sucrose than the other varieties, although the low amount of impurities is, to some extent, characteristic of the honey of the whole family (leguminosæ). The compositæ yield honey with about the average amount of organic non-sugars; the rosaceæ yield a product low in dextrin, but high in undetermined matter. Buckwheat and other polygonaceous honeys contain almost no sucrose, but give tests for tannins. Basswood honey is relatively high in dextrin, and that from poplar, oak, hickory and other trees, all of which contain considerable quantities of honeydew, are rich in both dextrin and ash. Pronounced tannin reactions are obtained in honey gathered from the flowers or plants of the sumac, hop and others rich in tannin. Tupelo, mangrove and sage honeys are distinguished by their high levulose content.

Browne found the average per cent of water in honey from the arid states of Arizona, Nevada, Utah, and Colorado was 15.60, and from the humid states of Minnesota, Wisconsin, Illinois, Missouri and Iowa was 18.88.

Hawaiian Honey.—This is characterized by its high ash and the presence of decided amounts of chlorides in the ash. Van Dine† states

^{*} U. S. Dept. of Agric., Bur. of Chem., Bul. 110 (1908).

[†] Ibid., p. 52.

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			Direct.		Invert	1		ರ	omplete	Complete Analysis				Tag G.
Source of Honey, according to Flower.	Number of Samples.	otaibemmI .O os ta	Constant	84° C.	3°°C.	87° C.	Water.	Invert Sugar.	Sucrose,	Ash.	Dextrin.	Undeter- mined.	Free Acid as Formic.	Reducing Su se Dextros
Alfalfa (Mediran satina)	œ	° V.	° V.	° V.	° V.	° V.	88	8	88	%	8	%	%	%
Maximum Minister (Teriolisms separe)	, ,	-17.5	1 20.5	+15.3	-24.5	+ 7.7	20.47	79.18	10.01	0.16	0.65	3.12	0.17	75.85 69.60
Maximum Minimum Ruckerpeat (Randowness fondowness)	?	- 3.2	- 18.2 - 6.0 +	7.4	-23.0 -12.9	$\frac{-23.0+13.8}{-12.9+5.1}$	20.24 14.54	78.15	0.00	0.00	2.46	7.45 3.11	0.10	74.86 67.37
Maximum Minimum Minimum Cotton (Corruption herbareum)	•	-14.7	- 17.4 - 16.2 +	8 8 4 0	-21.0 -19.8	+ 6.1	18.96	77.48	9.0	0.07	1.41	3.30	0.22	74.23 73.01
Maximum Minimum Basewood (Tilia th)	ر د	- 16.5 - 2.3	$\frac{-18.2 + 15.0}{-5.0 + 6.6}$	+15.0 +6.6	-22.0 - 7.5	+13.8 + 5.1	18.91	80.69	1.64	0.28	1.83	4.25 0.48	0.20	77.30
Maximum Minimum Sumac (Place)	, ; ; ;	- 9.3 + 3.7	- 13.7 - 0.3	-13.7 + 23.7 -0.3 + 9.7	- 16.8 - 1.3	+ 23.2	20.25	78.55 69.85	0.00	0.35	7.58	4.21 2.83	0.18	75.24 66.92
Maximum Minimum Orange (zirus auranisum) Turage (Nursea auranisum)	? Н (-11.1 - 4.6 -11.7	1 13.8 1 8.5 1 5.5	41 + + 6.88.	-17.6 +1 -11.4 + -19.3 +	-17.6 +14.4 -11.4 + 7.8 -19.3 + 6.6	19.25 18.17 16.99	73.73 68.61 77.57	0.36	0.90 0.08	6.42 1.66 0.45	6:68 3.89 4.31	0.18 0.09 0.08	70.63 65.73 74.31
Maximum Minimum Hawaiian honevdew and flowers	v	121.9	-24.6+ -23.4+	+ 6.2	-29.3 + -27.5 +	+ 4.6	18.38	72.40	4.36 1.65	0.08	2.69	6.02	0.06	69.36 69.06
Maximum Minimum	, ; ;	- 10.6 + 24.9	-15.1 +17.8		-18.2 +13.5	+34.8	17.80	76.55	5.27	0.48	10.01	4.18	0.15	73-33
Levorotatory honeys Maximum Minimum Destronolatory honeys	92	-21.9 + 3.7	- 24.8 - 0.3	+23.7	29-3 1-3	+23.2	26.88	83.36 10.01	10.01	0.03	7.58	7.45	0.25	79.86 59.61
Maximum		+ 24.9	$\mp +1$	+35.8 +28.5	+15.0	7.8 + 35.8 + 15.0 + 35.0 3.6 + 28.5 - 2.5 + 20.9	17.80	71.69	5.28 0.61	0.20	12.95	4-95 1.57	0.19	68.68 62.12

that the floral honey of Hawaii is largely from the blossoms of the algarroba (*Prosopis julifera*), while the honeydew honey, which, together with mixtures of honeydew and floral honey forms about two-thirds of the product of the Hawaiian Islands, comes largely from the exudations of the sugar-cane leaf-hopper (*Perkinsiella saccharicida*), and the sugar-cane aphis (*Aphis sacchari*). Honeydew honey is dextrorotatory, and for this reason has often been condemned as adulterated. It has a strong molasses-like odor, and often a very dark color. Bakers prefer it to algarroba honey, because of its baking and boiling properties.

The variation in the composition of Hawaiian honey is shown in the table on page 635, compiled from Browne's data.

Dextrorotarory Honey.—The U. S. standards define honey as levorotatory, thus excluding the larger part of the Hawaiian product, and also unimportant kinds of honey made from certain trees. Pure floral honey with no admixture of honeydew is seldom if ever dextrorotatory.

The following are the results obtained by Browne in the examination of detrorotatory honeys:

					н	awaiian.	
	Wild Penny- royal.	Poplar.	Hickory.	White Oak	Sugar Cane Honey- dew.	Honeydew and Flowers	Honeydew and Flowers.
Direct polarization at 20° C.* Invert polarization at 20° C Invert polarization at 87° C Water	+15.0	- 2.5 +20.9 17.02 65.80 3.10 0.76 10.19	+ 26.6 16.05 65.89 2.76 0.78 12.95 1.57	+28.6 13.56 65.87 4.31 0.79 10.49 4.98 0.08	15.46 64.84 5.27 1.29 10.01 3.13 0.15	16.29 67.81 2.57 1.02 9.65 2.66 0.14	66.85 2.41 0.80 8.62 3.52 0.13

^{*} Constant.

Adulteration of Honey.—The most common adulterants of honey are cane sugar, commercial invert sugar, and commercial glucose. Sometimes two or more adulterants are employed in the same sample. Gelatin is also said to be used. It appears to be a fact that bees may be made to feed upon cane syrup or commercial glucose, if these materials are placed in proximity to their hives, so that in some instances the adulterant

may be supplied through the medium of the bee. Sophisticated honey is often put up in tumblers or jars containing pieces of honeycomb, so that presence of the comb is by no means proof of its purity. Comb-honey, sold in the frame as sealed by the bees, is never adulterated, except when the bees are fed upon glucose or cane sugar.

Cane Sugar.—The following are typical analyses of honey adulterated with cane sugar:

	A.	В.	C.
Direct polarization	+34.7	+12	+ 1.2
Invert "	-24	-17.6	-21.5
Temperature	14°	·15°	19.5°
Sucrose (Clerget)	43.16%	21.8%	17.07%
Invert sugar	42.48%	60.03%	67.2%
Water	42.42%	21.15%	15.56%
Ash		.11%	0.06%

A strong right-handed polarization before inversion, coupled with a left-handed invert reading at 20°, is evidence of adulteration with cane sugar, or a product containing cane sugar.

Honey stored by bees fed on cane sugar is also characterized by its right-handed polarization. Although the bee inverts the larger part of the cane sugar in its body, this inversion is never as complete as in the case of nectar honey.

Glucose.—The following are typical analyses of honey adulterated with commercial glucose:

	A.*	B.	C.
Direct polarization	+ 147	+66.9	+101.5
Invert "	+135.2	+61.9	+ 99.0
Temperature	18°	20°	220
Sucrose (Clerget)	8.83%	3.76%	0.0%
Invert sugar	46.18%	74.66%	49.87%
Water	15.19%	21.40%†	23.7%
Ash	0.03%		

Care should be taken not to confuse honeydew honey with honey adulterated with glucose. Browne gives the following means of distinction:
(1) the difference in invert polarization between 20 and 87°, corrected to 77% invert sugar, (2) Beckman's iodine test (page 641), and (3) the

^{*} Both commercial glucose and added cane sugar.

König and Karsch test (page 642). He also finds the quantity and character of the ash, the acidity, and microscopic examination of value.

The following analyses of mixtures of commercial glucose and honey were made by A. H. Bryan.*

Mixture.			Invert Pol	rization—		Invert Sugar		Calculated Glucose.		
Glucose	Honey.	Constant Direct Polariza- tion at 20° C.		At 87° C.	Polariza- tion Differ- ence (87° –- 20°).	Before Inver- sion.	After Inver- sion.	Invert Polariza- tion at 87°+ 1.63.	Invert Polariza- tion at (20° C. + 17.5) + 1.93.	100- (Corrected Polarization Difference X 100+ 26.7)
%	%	° V.	° V.	° V.	° V.	%	%	%	%	%
100		+ 153.8	+153.34	+ 144.32		30.02	30.45	88.5	88.5	
50	50	+ 67.0	+ 65.67	+ 73.81	8.14	53.67	54.5c	45-3	43.1	56.9
20	80	+ 15.4			19.58	69.00	70.35	20.2	16.0	19.2
10	90	- 2.4	- 4.84	+ 18.59	23.43	74.42		11.4	6.6	8.8
5	95	- 11.5			25.96	75-74	77.8c		1.6	3.8
3 2	97	- 14.2				76.62			0.29	3.7
2	98	- 16.0	- 18.70	+ 8.14	26.84	76.64	78.34	5.0	0.00	1.2
1	99	- 18.2	- 20.90	+ 6.93	27.83	77.20		4.2	0.00	0.0
	100	- 19.5	- 22.11			77.68	78.9	3.2	0.00	0.0

Commercial Invert Sugar is the most difficult of detection of all the adulterants. Herzfeld's process† for the manufacture of invert sugar syrups consists in boiling for thirty to forty-five minutes 1 kilogram of refined sugar in 300 cc. of water with 1.1 gram of tartaric acid. Brown‡ gives the following analysis of the product made by this process:

Direct polarization at 20°	- 6.2
Constant polarization at 20°	- 9.5
Invert polarization at 20°	-16.9
Invert polarization at 87°	+ 4.8
Water	16.32%
Invert sugar	73-38%
Sucrose	4.36%
Ash	$\circ.\infty\%$
Dextrin	4.86%
	100.00
Acids as formic	0.06%

^{*}A. O. A. C. Proc., 1908, U. S. Dept. of Agric., Bur. of Chem., Bull. 122, p. 181.

[†] Zeits. ver. d. Zucker-Ind., 31, p. 1988.

^{\$} loc. cit., page 64.

This adulterant is best detected by Browne's test (page 642). Ley's test* has value in confirming the results of Browne's test, but should be used with caution, as American honeys do not react like the European.

Gelatin is indicated if a precipitate occurs in the diluted sample with a solution of tannic acid.

ANALYSIS OF HONEY.

Preparation of Sample.—In the case of strained honey, stir with a rod till any separated sugars are evenly distributed throughout the mass, or, if the honey has become solidified wholly or in part by crystallization, use a gentle heat on a closed water-bath to restore it to fluid form.

In the case of comb honey, cut with a knife across the top of the comb if sealed, and separate completely from the comb by straining through a 40-mesh sieve.

Determination of Moisture.†—Weigh 2 grams into a flat-bottom metal dish 2½ inches in diameter, which, together with 10 to 15 grams of fine quartz sand and a short stirring rod, has been previously tared, add 5 to 10 cc. of water, stir until the whole has been thoroughly incorporated, and dry to constant weight at 65 to 70° C. in a vacuum oven. Honeys of high purity usually dry in twelve hours, while those of the honeydew class rich in dextrin and gum require thirty-six hours, or longer.

Determination of Ash.—See page 614.

Polarization.—Direct and Invert at 20° C.—Proceed as directed under molasses (page 614), except that only alumina cream is used as a clarifier. To destroy birotation add a drop or two of ammonia before making up to the mark.‡

Invert at 87° C.—Invert a half normal portion in the usual manner in a 100-cc. flask, cool, add a few drops of phenolphthalein and enough sodium hydroxide to neutralize; discharge the pink color with a few drops of dilute hydrochloric acid, add from 5 to 10 cc. of alumina cream, make up to the mark and filter. Polarize in a 200-mm. tube at 87°, and multiply reading by 2.

Polarization at the temperature of 87° can most readily be effected by the use of a water-jacketed tube, as shown in Fig. 111. An all-metal tube, the interior of which is heavily gold-plated to avoid corrosion by acid, is preferable to one in which the inner tube is glass with a metal

^{*} Pharm. Zeits., 47, 1902, p. 603.

[†] Browne, U. S. Dept. of Agric., Bur. of Chem., Bul. 110, p. 18.

¹ Frühling, Zeits. öffentl. Chemie, 4, 1898, p. 410.

jacket, as in the latter leaky joints are liable to occur, due to uneven expansion. A tubulure is provided in the outer tube for a thermometer, so that the exact temperature may be noted. A tank of boiling water placed on a shelf above the polariscope is connected by rubber tubing with the jacketed tube as it rests in the polariscope, as shown in Fig. 111.

Determination of Reducing Sugars.—Determine by Allihn's method (page 608) in an aliquot of 25 cc. of a solution obtained by making 10 cc. of the solution prepared for polarization up to 250 cc. If desired the sugar may be determined by the volumetric Fehling process (page 591).

The reducing sugars may be calculated as dextrose as obtained from Allihn's table, or as levelose by multiplying the dextrose by 1.044.

Determination of Levulose.—Wiley's Method.*—This may be calculated approximately by the following formula:

$$l = \frac{100(1.0315A - a)}{(2.3919)26} = \frac{100(1.0315A - a)}{62.19},$$

in which l=levulose, a=the direct polarization at 20° of a solution of the normal quantity of honey made up to 100 cc. at 20°, and A=the direct polarization of the same solution at 87° C., 2.3919=the variation in polarization of 1 gram of levulose in 100 cc. of solution between 20 and 87° C., and 1.0315=the factor for converting the volume of the solution at 20° into that at 87° C.

Determination of Dextrose.†—Multiply the percentage of levulose as obtained in the preceding section by 0.915, thus obtaining the equivalent dextrose, and subtract this from the per cent of reducing sugars expressed as dextrose.

Determination of Sucrose.—Owing to the inaccuracies of Clerget's method as applied to honey, Browne recommends the following: Neutralize the free acid of 10 cc. of the solution used for invert polarization with sodium carbonate, make up to 250 cc. and determine the reducing sugars by Allihn's method. Subtract from the invert sugar thus obtained the invert sugar found before inversion, and multiply the difference by 0.95.

Determination of Dextrin.—Browne's Method.†—Weigh 8 grams of honey directly into a 100-cc. flask, add 4 cc. of water, and finally with

^{*} Principles and Practice of Agricultural Analysis, 1897, III, p. 267. Browne, loc. cit., p. 17.

[†] Browne, loc. cit., p. 17. Jour. Am. Chem. Soc., 28, 1906, p. 446.

continual agitation sufficient absolute alcohol to fill to the mark. Shake thoroughly and allow to stand twenty-four hours, or until the dextrin is deposited on the bottom and sides of the flask and the liquid is perfectly clear. Decant on a filter and wash the precipitate in the flask with 10 cc. of cold 95% alcohol, pouring the liquid finally on the filter. Dissolve the precipitate in the flask and on the filter in a little boiling, distilled water, collecting the solution in a tared platinum dish. Evaporate the liquid, and dry to constant weight at 100° C. If the alcohol precipitate is considerable, it should be dried at 70° C. in vacuo. After weighing, dissolve in water and make up to a definite volume according to the weight as follows:

Filter, determine invert sugar and sucrose in aliquots by copper reduction before and after inversion, and subtract the sum of these sugars from the total alcohol precipitate.

Determination of Acids.—Dissolve 10 grams of the honey in water and titrate with tenth-normal sodium hydroxide, using phenolphthalein as indicator. Express result as formic acid.

Beckman's Test for Glucose.*—Treat a mixture of equal parts of honey and water with a solution of iodine in potassium iodide. If glucose is present, a red or violet color (due to erythro- or amylo-dextrin) appears, the shade and intensity depending on the nature and amount of the glucose present.

Determination of Commercial Glucose in Honey.—Except for rough work, the method described on page 622 for calculating the per cent of commercial glucose from the sucrose and from the direct polarization is not recommended for use with honey and other products wherein the invert sugar is so large as to considerably affect its accuracy. In this case, it is best after inversion to polarize the sample at 87° C., the temperature at which the reading due to invert sugar would theoretically be o. At this temperature, any considerable right-handed polarization can be accounted as due to commercial glucose. (See page 639.)

As in the case of molasses, the writer advocates assuming 175° as the direct polarization of the glucose used, this being about the maximum reading for a normal solution of 42°-Bé. glucose. Lythgoe has shown

^{*} Zeits. Anal. Chem., 35, 1896, p. 267.

that in polarizing at high temperatures samples of saccharine products containing commercial glucose, certain precautions have to be observed not necessary when cane or invert sugar are the only sugars present. Thus, a normal solution of glucose, when polarized at 87° C., has a lower reading than in the cold, the difference being doubtless due partly at least to the expansion of the liquid. Again, on subjecting a normal solution of glucose to inversion with acid, as in Clerget's process, and heating to 87° C., it will be found impossible to get a constant reading, but the reading will drop rapidly, due to a partial hydrolysis of the maltose or dextrin.

In honey and other preparations containing much invert sugar and commercial glucose, it is best to proceed as follows: Divide the polarization at 87° by 163* and multiply the result by 100 for the percentage of commercial glucose in terms of glucose polarizing at 175°. It should be borne in mind that the results by even this method are only approximate, as genuine honey is more or less dextrorotatory at 87° C.

The following formula is used by European chemists: $G = \frac{b+17.5}{1.93}$ in which G = the per cent of commercial glucose, and b = the polarization after inversion at 20° C.

Browne's Test for Commercial Invert Sugar.†—Reagent.—This should be freshly prepared each time before using. Shake 5 cc. of c. p. anilin with 5 cc. of water, and add sufficient glacial acetic acid (2 cc.) to just clear the emulsion.

Process.—Treat 5 cc. of a 1:1 solution of the honey in a test tube with 1 to 2 cc. of the anilin reagent, allowing the latter to flow down the walls of the tube so as to form a layer upon the honey solution. If, when the tube is gently agitated, a red ring forms beneath the anilin solution, this color becoming gradually imparted to the whole layer, artificial invert sugar is present. This reaction is due to furfural formed during the high temperature employed in the commercial processes of inversion. Boiling genuine honey also causes the formation of furfural, but this treatment impairs the flavor and is probably never practiced.

Distinction of Honeydew and Glucose Honeys.—Method of König and Karsch.‡—Dissolve 40 grams of honey in a cylinder in water, and

^{*}The true polarization at 87° C. of a normal solution of glucose subjected to inversion and neutralization as above (but without the use of the clarifier), will be about 93% that of the direct polarization of the sample in the cold. Hence $175 \times 0.93 = 162.7$.

[†] U. S. Dept. of Agric., Bur. of Chem., Bul. 110, p. 68.

[‡] Zeits. anal. Chem., 34, 1895, p. 1. U. S. Dept. of Agric., Bur. of Chem., Bul. 110, p. 63.

make up to 40 cc. Transfer 20 cc. of the homogeneous solution to a 250-cc. flask and fill to mark with absolute alcohol with slow addition and constant shaking, and then allow to stand two or three days, with occasional agitation. At the end of this time all the dextrin has settled out. After shaking the solution, filter and evaporate 100 cc. of the filtrate until free from alcohol. To the liquid residue add a little subacetate of lead and sodium sulphate, make up to 20 cc. with water, and polarize the filtered solution. Dextrorotatory natural honeys show by this method a levorotation; honeys adulterated with dextrose or glucose to the extent of 25% or more, a dextrorotation. In case the honey contains a large amount of sucrose, the solution should be inverted with hydrochloric acid before polarizing.

BEESWAX.—The purity of beeswax is best established by determining its melting-point, its specific gravity, its saponification equivalent, and its refractometric reading. The melting-point of pure wax is about 64° C., while that of paraffin, its chief adulterant, is from 52 to 55° C. Its saponification equivalent should be from 87.8 to 107, while that of paraffin is o.

Method of Determining Specific Gravity of Beeswax.*—Place a weighed rod of the wax, about 1 to 1.5 cm. long by 0.5 cm. diameter, in an accurately marked 50-cc. flask, and run in water from a burette till the water level reaches the mark. 50 cc. minus the burette reading represent the volume occupied by the wax. The rod should be made to lie flat on the bottom of the flask, so that the incoming water will force its end against the sides and prevent the end from rising above the mark. The weight of the rod, divided by its volume gives its specific gravity. The specific gravity of various mixtures of wax of 0.969 specific gravity and paraffin of 0.871 are given in the following table, prepared by Wagner, so that from the specific gravity of the mixture the percentage of paraffin can be calculated:

Wax	Paraffin	Specific	Wax	Paraffin	Specific
(Percentage).	(Percentage).	Gravity.	(Percentage).	(Percentage).	Gravity.
25 50	100 75 50	.871 .893 .920	75 80 100	25 20 —	-942 -948 -969

The Refractometer Reading is most useful in establishing the purity of wax. Observations with this instrument are best made at 65° and

^{*} U. S. Dept. of Agric., Div. of Chem., Bul. 13, p. 842.

great care should be taken in the case of the Zeiss butyro-refractometer not to exceed this temperature, or injury to the instrument may result.

The Abbé refractometer may be used with perfect safety and, when available, is to be preferred for the examination of beeswax. Many food laboratories are, however, not equipped with the Abbé, but nearly all find the butyro-refractometer indispensable. The latter instrument was primarily designed for such substances as butter and lard, so that the

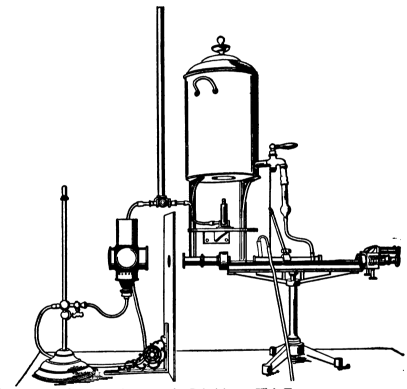


Fig. 111.—Apparatus for Polarizing at High Temperatures.

manufacturers did not intend it to be subjected to as high a temperature as 65°. They have, however, assured the author that if care be taken to bring the temperature very slowly and gradually to the required degree, 65°, and to avoid also sudden cooling, the cement that secures the prisms in place will not be appreciably affected; otherwise cracking or loosening of the cement would be liable to occur after a time.

At 65° C. pure beeswax should have a reading on the butyro-refractometer of 30 to 31.5,* while that of paraffin is from 11 to 14.5.†

CONFECTIONERY.

The composition of confectionery is more complex than that of the saccharine products hitherto considered. As a rule, cane sugar, or one of its products, as molasses, forms the basis of most of the confections. Commercial glucose is also a common ingredient, while a large variety of such materials as eggs, butter, chocolate, various flavoring extracts, spices, nuts, and fruits, enter into the composition of confectionery.

U. S. Standard Candy is candy containing no terra alba, barytes, talc, chrome yellow, or other mineral substances or poisonous colors or flavors, or other ingredients injurious to health.

Adulteration.—Of late the adulteration of confectionery has been held largely in check by the National Confectioners' Association of the United States, which has fixed high standards of purity, and has been very zealous in restricting the use of harmful adulterants.

Commercial glucose is not regarded as an adulterant of confectionery by the above-named association and by but few of the state laws. On the contrary, any ingredient, other than color, that has no food value, may logically be considered as an adulterant. Under this head are included such substances as paraffin, as well as make-weight mineral matters, such as terra alba, talc, or calcium sulphate.

Most of the actually harmful ingredients employed in confectionery have been inherent in the coloring matters, or in the alcohol or fusel oil used in the manufacture of brandy drops and allied confections.

Colors in Confectionery.—A very wide range of colors is necessarily employed in the manufacture of confectionery, and the almost endless variety of coal-tar dyes now available lend themselves most readily to the confectioner's needs. Elsewhere, under "colors," lists of injurious and non-injurious dyes are given as compiled by the National Confectioners' Association, though it is not always readily apparent how the lines are drawn.

The tinctorial power of these dyes is so high that the actual amount of substance contained in a thin coating of the color on the outside of the candy is exceedingly small, so that it is doubtful whether serious cases of injury have ever arisen from their use.

^{*} nD, 1.4452 tb 1,4463.

Such was not the case formerly, before the prevalence of the coal-tar dyes, when such poisonous mineral pigments as chromate of lead were frequently used. Only one or two instances of the use of lead chromate in candy have come to the author's attention within ten years, since more satisfactory and harmless yellow colors among the azo-dyes are now obtainable.

ANALYSIS OF CONFECTIONERY.—The following have been submitted by the author as provisional methods of procedure for the A. O. A. C.:*

(1) Products of Practically Uniform Composition Throughout.—
(a) Lozenges and Other Pulverizable Products.—Grind in a mortar or mill to a fine powder. For total solids, weigh from 2 to 5 grams of the powdered sample in a tared platinum dish, and dry in a McGill oven to constant weight.

For Ash, ignite the residue from total solids in the original dish, observing the precautions given under sugar (p. 586), and molasses (p. 624).

- (b) Semi-plastic, Syrupy, or Pasty Products.—Weigh 50 grams of the sample into a 250-cc. graduated flask, mix thoroughly or dissolve, if soluble, in water, and fill to the mark. Be sure that the solution is uniform, or, if insoluble material is present, that it is evenly mixed by shaking before taking aliquot parts for the various determinations. For total solids and ash, measure 25 cc. of the above solution or mixture into a tared platinum dish, and proceed as directed under (a).
- (2) Confectionery in Layers or Sections of Different Composition.—When it is desired to examine the different portions separately, they should be separated mechanically with a knife, when possible, and treated as directed under (1).
- (3) Sugar-coated Fruit, Nuts, etc.—In case of a saccharine coating inclosing fruit, nuts, or any less readily soluble material, dissolve or wash off the exterior coating in water, which may, if desired, be evaporated to dryness for weighing, and proceed as in (1).
- (4) Candied or Sugared Fruits.—Proceed as in the examination of fruits (Chapter XXI).

Detection of Mineral Adulterants.—As in the case of molasses, a considerable quantity, say 100 grams, should be reduced to an ash for examination for mineral adulterants, such as talc, calcium sulphate, and iron oxide, which are detected by regular qualitative tests.

^{*} U. S. Dept. of Agric., Bur. of Chem., Bul. 65, p. 44.

Detection of Lead Chromate.—Fuse the ash in a porcelain crucible with a mixture of sodium carbonate and potassium chlorate, boil the fused residue with water, neutralize with acetic acid, filter, and treat the filtrate with barium chloride or lead acetate solution. A yellow precipitate indicates a chromate. Treat the insoluble part of the fusion with nitric acid, and test for lead in the usual manner.

If a drop of ammonium sulphide be applied to a piece of confectionery colored with lead chromate, it will produce a black coloration.

Determination of Ether Extract.—The ether extract includes the fat derived from chocolate, eggs, or butter, as well as any paraffin present. Measure 25 cc. of the 20% solution (1) (b) (p. 646) into a very thin, readily frangible glass evaporating-shell (Hoffmeister's Schälchen), containing 5 to 7 grams of freshly ignited asbestos fiber; or, if impossible to thus obtain a uniform sample, weigh out 5 grams of the mixed, finely divided sample into a dish, and wash with water into the asbestos in the evaporating-shell, using, if necessary, a small portion of the asbestos fiber on a stirring-rod to transfer the last traces of the sample from dish to shell. Dry to constant weight at 100°, after which cool, wrap loosely in smooth paper, and crush into rather small fragments between the fingers, carefully transferring the pieces with the aid of a camel's-hair brush to an extraction-tube, or to a Schleicher and Schull cartridge for fat extraction. Extract with anhydrous ether or with petroleum ether in a continuous extraction apparatus for at least twenty-five hours. fer the solution to a tared flask, evaporate the ether, dry in an oven at 700° C. to constant weight, and weigh.

Determination of Paraffin.—Add to the ether extract in the flask, as above obtained, 10 cc. of 95% alcohol, and 2 cc. of 1:1 sodium hydroxide solution, connect the flask with a reflux condenser, and heat for an hour on the water-bath or until saponification is complete. Remove the condenser, and allow the flask to remain on the bath till the alcohol is evaporated off, and a dry residue is left. Treat the residue with about 40 cc. of water, and heat on the bath, with frequent shaking, till everything soluble is in solution. Wash into a separatory funnel, cool, and extract with four successive portions of petroleum ether, which are collected in a tared flask or capsule. Remove the petroleum ether by evaporation, and dry in the oven to constant weight.

It should be noted that any phytosterol or cholesterol present in the fat would come down with the paraffin, but the amount would be so insignificant that, except in the most exacting work, it may be disregarded.

The character of the final residue should, however, be confirmed by determining its melting-point and specific gravity, and by subjecting it to examination in the butyro-refractometer. The melting-point of paraffin is about 54.5° C.; its specific gravity at 15.5° C. is from 0.868 to 0.915, and on the butyro-refractometer the reading at 65° C. is from 11 to 14.5.

Determination of Starch.—Measure gradually 25 cc. of a 20% aqueous solution or uniform mixture of the sample into a hardened filter or Gooch crucible, or transfer by washing 5 grams of the finely powdered substance to the filter or Gooch, and allow the residue on the filter to become airdried. Extract with five successive portions of 10 cc. of ether, then wash with 150 cc. of 10% alcohol, and finally with 20 cc. of strong alcohol. Transfer the residue to a large flask and boil gently for four hours with 200 cc. of water and 20 cc. of hydrochloric acid (specific gravity 1.125), the flask being provided with a reflux condenser. Cool, neutralize with sodium hydroxide, add 5 cc. of alumina cream, and make up the volume to 250 cc. with water. Filter and determine the dextrose in an aliquot part of the filtrate by any of the various Fehling methods. The weight of the dextrose multiplied by 0.0 gives the weight of the starch.

Polarization of Confectionery.—As a clarifier use either alumina cream or subacetate of lead, according to the nature and opacity of the sample. Ordinarily alumina cream is best, but in dark-colored samples, or those in which molasses has been used, it is sometimes necessary to employ the subacetate. When starch is absent, and the sample is practically soluble, polarize and invert in the usual manner (p. 588). Where considerable starch or insoluble matter is present, use the double-dilution method of Wiley and Ewell (p. 620), thus making due allowance for the volume of the precipitate.

Cane sugar, invert sugar, and dextrin, are determined as directed for honey.

Commercial glucose is roughly determined by polarizing the sample at 87° C., as in the case of honey (p. 639).

Confectionery is made in such a wide variety of forms, and these differ in consistency to such an extent that commercial glucose of all available degrees of density can be utilized to advantage in one product or another. In this respect confectionery is unlike honey and molasses, wherein a fairly uniform grade of commercial glucose is necessarily used for mixing, this grade being naturally selected with reference to its similarity in density to the molasses. On this account the glucose factor used

for honey and molasses (175) may in some varieties of confectionery be too high.

Determination of Alcohol in Syrups Used in Confectionery.—(Brandydrops.)—Open each drop by cutting off a section with a sharp knife, and collect in a beaker the syrup of from 15 to 25 of the drops, which will usually yield from 30 to 50 grams of syrup. Strain the syrup into a tared beaker through a perforated porcelain filter-plate in a funnel to separate from particles of the inclosing shell, and ascertain the weight of the syrup. Wash into a distilling-flask, dilute with half its volume of water, and distil off into a tared receiving-flask a volume equal to the original volume of syrup taken. Ascertain the weight of the distillate and its specific gravity by means of a pycnometer. Multiply the percentage by weight of alcohol corresponding to the specific gravity, as found in the tables on page 661 et seq., by the weight of the distillate, and divide this by the weight of syrup taken. The result is the per cent by weight of alcohol in the syrup.

Detection of Colors.—It is sometimes necessary to macerate a considerable mass of the material to remove the color, which is, however, in the majority of cases readily soluble. The insoluble colors are nearly all mineral pigments to be looked for in the ash, as in the case of chromate of lead (p. 647). Frequently the coloring matter is confined to a thin outer layer, which is readily washed off.

The solution of the dyestuff is examined as directed under colors.

Detection of Arsenic.—Arsenic may be present through impure coloring-matter. If the color is confined to an exterior coating, this should be washed off and examined. If distributed through the mass, a solution of the whole should be taken. Examine for arsenic by the Gutzeit or Marsh method, as directed under glucose (p. 632).

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CHAPTER XV.

ALCOHOLIC BEVERAGES.

Alcoholic Fermentation.—In a broad sense all alcoholic liquors are saccharine products, in that they are essentially the result of the fermentation of sugar. In the case of fruits, the sugar already exists as such in their juices, which, when expressed, almost immediately on exposure to the air begin to undergo spontaneously the process of alcoholic fermentation, in accordance with the reaction:

(1)
$$C_0H_{12}O_0=2C_2H_0O+2CO_2$$
.

Describes or Alcohol Carbon dioxide

In the case of grains the process is more complex, involving a preliminary saccharous fermentation, whereby the starch is first transformed into sugar.

Thus

(2)
$${}_{2}C_{e}H_{10}O_{5} + H_{2}O = C_{e}H_{10}O_{5} + C_{e}H_{12}O_{e}$$

Starch Dextrose

(3)
$$C_6H_{10}O_5+H_2O=C_6H_{12}O_6$$

Dextrope

The process of alcoholic or vinous fermentation is largely dependent upon the presence of various species of yeasts, which either exist from the first in the expressed juices themselves, as in the case of wines, being derived from the skins of the grapes and from the air, or are introduced with some degree of selection, as in the case of beer.

In the juices of most fruits the sugar exists in the form of sucrose, mixed with variable amounts of invert sugar resulting from the inversion of the sucrose due to the action of ferments, such as invertase, a soluble ferment of yeast. The invert sugar nearly always predominates, and in some juices, as for instance the grape, nearly all the sugar has been inverted.

The above reaction, No. 1, illustrating the splitting up of grape sugar into alcohol and carbon dioxide, does not represent the practical yield of alcohol under ordinary conditions that occur in vinous fermentation, for, as a matter of fact, instead of 51.11 parts of alcohol and 48.89 parts carbon dioxide, which would theoretically result as above from the fermentation of 100 parts of dextrose, only about 95% of the theoretical yield can be obtained, so that in practice it is possible to form but about 48.5% alcohol and 46.5% carbon dioxide. The balance, amounting to some 5%, consists chiefly of glycerin, succinic acid, and traces of various compounds, including some of the higher-boiling alcohols (propyl, butyl, and amyl) and their ethers, which form the fusel oil of the distilled liquors.

Vinous fermentation takes place most readily in slightly acid liquids, at a temperature ranging from 25° to 30° C.

It is convenient to divide alcoholic beverages into two main groups, first the fermented and second the distilled liquors. The fermented liquors naturally subdivide themselves into (a) the products of the direct spontaneous fermentation of saccharine fruit juices, such, for example, as those of the apple and the grape, to form cider and wine respectively, and (b) the malted and brewed liquors, such as beer and ale, produced by the conversion of the starch of grain into sugar, and the final alcoholic fermentation of the latter.

The distilled liquors include such products as whiskey, brandy, rum, and gin, wherein alcoholic infusions prepared by previous fermentation in various ways are further subjected to distillation.

Alcoholic Liquors and State (or Municipal) Control.—The mere adulteration of liquors does not constitute the only feature which brings them within the scope of the public analyst's work and renders them especially amenable to stringent laws. Indeed, it is often a far more important question for the analyst to decide by his results whether or not the samples submitted to him, by police seizure or otherwise, are sold in violation of the regulations in force in his particular locality governing the liquor traffic.

A common regulation in no-license localities fixes the maximum per cent of alcohol which shall decide whether or not a liquor is legally a temperance drink, and can be sold as such with impunity. From its low content in alcohol, an analyst's findings regarding a certain sample may exonerate the dealer suspected of violating this law, while yet by the very reason of its being low in alcohol the same sample would be placed

in the adulterated list as regards non-conformance to a standard of purity. While the raising of revenue is one purpose for the existence of these laws bearing on liquor license, an equally important object sought to be gained is doubtless the repression of intemperance.

Toxic Effects.—A popular impression seems to exist that the toxic effects of an adulterated liquor are far worse from a temperance stand-point than those of a sample of good standard quality, and it is a common experience of the public analyst to have submitted to him by well-meaning temperance advocates samples which are alleged to have caused the worst forms of intoxication, and are thus suspected of being impure. As a matter of fact the chief adulterants of liquors are water, sugar, and, in the case of beer, various bitter principles and vegetable extractives, none of which are on record as being in themselves actively toxic.*

Alcohol is the one ingredient of liquor which, more than any other, produces a marked physiological effect. Many liquors, especially those of the distilled variety classed as adulterated, are so considered by reason of their low alcoholic content through watering or otherwise, hence this commonest form of adulteration, far from being detrimental in itself, is actually helpful to the temperance cause.

Details of Liquor Inspection.—The same precautions should be carefully observed by officers making seizures of liquors for analysis, as by food inspectors, regarding safe delivery of the samples to the analyst. The following instructions are circulated by the State Board of Health of Massachusetts, which has in charge the inspection of liquors, concerning the taking of samples in that state and the transmission to the analyst:

DIRECTIONS FOR TAKING SAMPLES FOR ANALYSES.

The officer making a seizure, or taking samples of beer, should note at the time of such seizure the general appearance of the liquor,—as to whether it is clear or cloudy, whether it is still or has a strong head.

If the liquor is in bottles, take at least one pint bottle; if in barrels, draw a pint bottle from each. Request the owner to seal each sample taken. If the bottles have cork stoppers, cut the stoppers off level with the top of the bottle and cover with wax; if with patent stoppers, a little wax placed upon the wire at the point where it lays against the neck of the bottle is sufficient. If the owner refuses to seal it, then the officer

^{*} The writer refers to substances intentionally added, and not to accidental impurities, such as arsenic, etc., that are occasionally found.

should seal it in his presence, calling his attention to the fact. Before leaving the premises, place upon the bottle a label or tag, with the date, the name of the owner, and the name of the officer upon it, and also the name of the town or city. Then place in a box, with the certificate required by law, and forward without delay to the analyst.

FORM OF LABEL.

Date of seizure Owner Kind of liquor	19
first part of which is filled out a part, containing the data of analy and returned by him to the off	is a certificate like the following, the nd signed by the officer, while the second rsis, is filled out and signed by the analysticer. Such a certificate is nearly always without the personal appearance of the
To the State Board of Health: I send herewith a sample of	19
Ascertain the percentage of	alcohol it contains, by volume, at sixty to me a certificate herewith upon the
	Officer.
COMMONWEALT	H OF MASSACHUSETTS.
No	STATE BOARD OF HEALTH. BOSTON,
	received by me isper cent of alcohol, enheit.
SEAL.	Analyst State Board of Health.

A convenient method for recording analyses is by the employment of numbered library cards, which bear the same number as the certificates and are kept by the analyst.

The following is a convenient form:

No	Analyzed. Wt. flask and alc. Wt. flask. Wt. alc. Sp. gr. alc. (60°) Per cent alcohol. Reported.
Received	
Sealed.	
Condition	
Kind of bottle	
Registered	

METHODS OF ANALYSIS COMMON TO ALL LIQUORS.

Specific Gravity.—This should be taken at 15.6° or calculated to that temperature. The most convenient mode of procedure is to bring the temperature of the sample somewhat below that point by allowing the flask containing it to stand in cold water, and to have everything in readiness to make the determination when 15.6° temperature has been reached, either by the hydrometer spindle in a glass cylinder, by the Westphal balance, or by the pycnometer. The latter is by far the most accurate, especially if it is of the form which is fitted with a the momenter-stopper.

Detection of Alcohol.—It is rarely necessary to make a qualitative test for alcohol in liquors, since it is almost invariably present even in many of the so-called temperance drinks, at least in small amount. Indeed in many localities a beverage is legally a temperance drink that contains not more than 1% alcohol by volume.

The Iodojorm Test.—Alcohol, when present in aqueous solution to the extent of 0.1% or more, may be detected by the iodoform test. The solution is warmed in a test-tube with a few drops of a strong solution of iodine in potassium iodide, after which enough sodium hydroxide solution is added to nearly decolorize. On standing for some time a yellow precipitate of iodoform will appear if alcohol be present, or at once if there is a considerable amount, and the characteristic odor of iodoform will be rendered apparent, even when the precipitate is so slight as to be almost imperceptible. This iodoform precipitate is crystalline, showing under the microscope as star-shaped groups or hexagonal tablets.

It should not be forgotten that other substances than alcohol give the reaction, as lactic acid, acetone, and various aldehydes and ketones.

Pure methyl or amyl alcohol or acetic acid do not thus react.

Berthelot recommends benzoyl chloride as a reagent for detecting alcohol. By warming a mixture of a few drops of benzoyl chloride with the solution to be tested, and adding a little sodium hydroxide, ethyl benzoate is formed, recognizable by its distinctive odor. This reaction is delicate to 0.1% alcohol. The presence of other alcohols than ethyl produces ethers of characteristic odor.

Hardy's Test for Alcohol consists in shaking the aqueous solution with some powdered guaiacum resin, filtering, and adding to the filtrate a little hydrocyanic acid and a drop of dilute copper sulphate solution. A blue coloration considerably deeper than that due to the copper salt is indicative of alcohol.

Methyl Alcohol in spirits is tested for as described on pp. 749-752.

Determination of Alcohol.—In the case of carbonated liquids it is necessary to first expel the free carbon dioxide, which is readily accomplished by pouring the liquor back and forth from one beaker to another, from time to time removing the excess of froth from the top of the vessel by the aid of the hand. Or, the sample may be shaken vigorously in a large separatory funnel, and the still liquor drawn off from below the froth, repeating the operation several times if necessary. In either case the mechanical treatment should be continued till the liquor is comparatively quiet and free from foam.

(1) By Distillation.—This is by far the most accurate method of determining alcohol, and should be carried out in all cases where any legal controversy is apt to be involved. Into a flask of 250 to 400 cc. capacity introduce a convenient quantity of the liquor, which should be accurately weighed or measured, according to whether the percentage by weight or measure is desired. The following are suitable quantities: Distilled liquors, 25 grams or cc.; cordials, 25 to 50 grams or cc.; wines, ciders, and malt liquors, 100 grams or cc. In the case of wines or ciders which have undergone acetic fermentation, add 0.1 to 0.2 gram of precipitated calcium carbonate or neutralize with standard alkali.

Dilute the liquid to 150 cc. and distil into a 100-cc. flask. Nearly all alcoholic liquors, if comparatively free from carbon dioxide, will boil without undue frothing or foaming. New wine will occasionally give trouble in this regard, but foaming may usually be prevented in this

no-

case by the addition of tannic acid. In case of wine, cider, and beer all the alcohol will have passed over in the first 75 cc. of the distillate. or three-fourths the original measured volume, but with distilled liquors high in alcohol the process had better be continued till nearly 100 cc. or the original volume taken have passed over. If the condenser is of glass. one can observe when all the alcohol has been distilled over, for the reason that the mixed alcohol and water vapors in the upper portion of the condenser present a striated or wavy appearance, readily apparent so long as the alcohol is passing over, while after all the alcohol has been distilled. the condenser-tube appears perfectly clear. The distillation is thus continued for some time after this striated appearance has ceased. The distillate in the receiving glass is finally made up to the mark or to the original volume of the liquor taken. Strictly speaking, the measurements before and after distillation should be made at 15.6° C., but, excepting in case of distilled liquors, no appreciable error results from making both measurements at the same or room temperature. Another precaution formerly thought necessary was to have the delivery-tube from the condenser pass below the level of a little water in the receiving-flask from the start, but equally accurate results have been obtained by simply allowing the end of the condenser-tube to enter the narrow-necked flask.

Fig. 112 shows a bank of six stills of the kind used in the author's laboratory for alcohol determination in liquors. In each still the vertical glass worm-condenser, the round-bottomed distilling-flask, and the lamp, are supported by rings held by a single upright rod. The receiving-flask is readily connected with the condenser by means of a single bent tube provided with a rubber stopper. The cold-water pipe supplying the condensers is shown at the top, and the gas-supply pipe at the bottom.

The distillate, made up to 100 cc., is thoroughly shaken and its specific gravity taken at exactly 15.6° in a pycnometer, or by the Westphal balance. From the specific gravity the corresponding percentage of alcohol by weight or volume, or the grams per 100 cc. in the distillate, is ascertained by reference to the accompanying tables.

To obtain percentage of alcohol by weight in the sample, multiply the per cent by weight in the distillate by the weight of the distillate, and divide by the weight of the sample taken; to obtain per cent by volume, multiply the per cent by volume in the distillate by 100, and divide by the volume of the sample used.

(2) From the Specific Gravity of the Sample.—In the case of distilled liquors having very little residue, an approximation to the true

percentage of alcohol may be obtained by using the alcohol table in connection with the specific gravity of the liquor itself. The accuracy of this method depends largely on the freedom from residue, being absolutely correct for mixtures of alcohol and water only.

(3) By Evaporation.—Determine the specific gravity of the sample, evaporate a measured portion of the liquor (50 or 100 cc.) in a porcelain

Fig. 112.—Bank of Stills for Alcohol Determination.

dish over the water-bath to one-fourth its bulk, make up to its original volume with distilled water, and determine the specific gravity of this second or dealcoholized portion. Add I to the original specific gravity, and from this subtract the second specific gravity. The difference is the specific gravity corresponding to the alcohol in the liquor, the percent of which is found from the table.

Example.—Suppose the specific gravity of the original sample to be 0.9900 while that of the dealcoholized sample is 1.0009. Then 1.9900—1.0009=0.9891. ... Per Cent by volume of alcohol=8.10.

SPECIFIC GRAVITY AND PERCENTAGE OF ALCOHOL. (According to Hehner.)

0	Abso	lute Alc	oʻhol.		Abso	lute Alc	ohol.		Abso	lute Alc	ohol.
Spec. Grav. at 15.6° C.	Per Cent by Weight	Per Cent by Vol- ume.	Grams per 100 cc.	Spec. Grav. at 15.6° C.	Per Cent by Weight	Per Cent by Vol- ume.	Grams per 100 cc.	Spec. Grav. at 15.6° C.	Per Cent by Weight	Per Cent by Vol- ume.	Grams per 100 cc.
1.0000	0.00	0.00	0.00								
0.9999 8 7 6 5 4 3 3 2 1 0 0 0.9989 8 7 6 5 4 3 2 1 0 0	0.05 0.11 0.16 0.21 0.26 0.32 0.37 0.42 0.47 0.53 0.68 0.74 0.79 0.84 0.89 0.95 1.06	0.07 0.13 0.26 0.33 0.40 0.45 0.53 0.60 0.66 0.73 0.93 0.93 0.93 1.10 1.13 1.19 1.26	0.05 0.11 0.16 0.21 0.26 0.32 0.37 0.47 0.53 0.68 0.74 0.74 0.89 0.95 1.00	0.9959 8 7 6 5 4 3 2 1 0 0.9949 8 7 6 5 4 3 2 1 0	2.33 2.39 2.44 2.50 2.56 2.67 2.72 2.78 2.83 2.89 2.94 3.06 3.12 3.18 3.24 3.29 3.35 3.41	2.93 3.00 3.07 3.14 3.28 3.35 3.42 3.49 3.55 3.62 3.69 3.76 3.83 3.90 3.98 4.05 4.12 4.20	2.32 2.38 2.43 2.45 2.55 2.60 2.65 2.70 2.81 2.87 2.92 2.98 3.10 3.16 3.22 3.23 3.33 3.39	0.9919 8 7 6 5 4 3 2 1 0 0 0.9909 8 7 6 5 4 3 2 1 0	4.69 4.75 4.81 4.87 4.90 5.06 5.12 5.25 5.37 5.37 5.56 5.62 5.62 5.75 5.87	5.86 5.94 6.02 6.10 6.24 6.32 6.40 6.48 6.55 6.63 6.71 6.78 6.86 6.94 7.01 7.09 7.17 7.25 7.32	4-65 4-71 4-77 4-83 4-95 5-07 5-14 5-20 5-32 5-32 5-35 5-57 5-57 5-57 5-58 5-76
0.9979 8 7 6 5 4 3 2 1 0	1.12 1.19 1.25 1.31 1.37 1.44 1.50 1.56 1.62 1.69	1.42 1.49 1.57 1.65 1.73 1.81 1.88 1.96 2.04 2.12	1.12 1.19 1.25 1.31 1.37 1.44 1.50 1.56 1.61 1.68	0.9939 8 7 6 5 4 3 2 1 0	3-47 3-53 3-59 3-65 3-71 3-76 3-82 3-88 3-94 4-00	4-34 4-42 4-49 4-56 4-63 4-71 4-78 4-85 4-93 5-08 5-08	3-45 3-51 3-57 3-63 3-69 3-74 3-80 3-85 3-91 3-97 4-03 4-09	0.9899 8 7 6 5 4 3 2 1 0	5.94 6.00 6.07 6.14 6.21 6.28 6.36 6.43 6.50 6.57 6.64 6.71	7-40 7-48 7-57 7-66 7-74 7-83 7-92 8-01 8.10 8.18	5.88 5.94 6.01 6.07 6.14 6.21 6.29 6.36 6.43 6.50
7 6 5 4 3 2 1	1.87 1.94 2.00 2.06 2.11 2.17 2.22 2.28	2-35 2-43 2-51 2-58 2-62 2-72 2-79 2-86	1.86 1.93 1.99 2.05 2.10 2.16 2.21 2.27	7 6 5 4 3 2 1	4-19 4-25 4-31 4-37 4-44 4-50 4-56 4-62	5-24 5-32 5-39 5-47 5-55 5-63 5-71 5-78	4.16 4.22 4.28 4.34 4.40 4.46 4.52 4.58	7 6 5 4 3 2 1	6.78 6.86 6.93 7.00 7.07 7.13 7 20 7.27	8.45 8.54 8.63 8.72 8.80 8.88 8.96 9.04	6.70 6.78 6.85 6.92 6.99 7.05 7.12 7-19

_	Abs	olute Alc	ohol.	 	Abs	olute Ale	ohol.		Absc	olute Alc	ohol.
Spec. Grav. at 15.6° C.	Per Cent by Weight	Per Cent by Vol- ume.	Grams per 100 cc.	Spec. Grav. at 15.6° C.	Per Cent by Weight	Per Cent by Vol- ume.	Grams per 100 cc.	Spec. Grav. at 15.6° C.	Per Cent by Weight	Per Cent by Vol- ume.	Grams per 100 cc.
0.9 879 8 7 6	7-33 7-40 7-47 7-53	9.13 9.21 9.29 9.37	7-24 7-31 7-37 7-43	0.9829 8 7 6	11.00	13-71 13.81	10.73 10.81 10.89 10.95	0.9779 8 7 6	14.91 15.00 15.08 15.17	18.68	14.66 14.74 14.83
5 4 3 2 1	7.80	9-45 9-54 9-62 9-70 9-78	7.50 7.57 7.63 7.70 7.77	5 4 3 2 1	11.38 11.46 11.54	13.99 14.09 14.18 14.27	11.03 11.11 11.18 11.26 11.33	5 4 3 2 1	15.25 15.33 15.42 15.50 15.58	18.78 18.88 18.98 19.08	14.98 15.07 15.14 15.21
0.986 9 8	8.00 8.07 8.14	9.86 9.95 10.03	7.89 7.96 8.04	0.9819	11.69 11.77 11.85	14.46 14.56 14.65	11.56	0.9769 8 7 6	15.75 15.83	19.39 19.49 19.59	15.38 15.46 15.54
6 5 4 3 2	8.29 8.36 8.43 8.50	10.21 10.30 10.38 10.47 10.56	8.10 8.17 8.24 8.31 8.38 8.45	5 4 3 2	12.00 12.08 12.15 12.23	14.84 14.93 15.02 15.12	11.70 11.78 11.85 11.92 12.00 12.08	5 4 3 2	16.00 16.08 16.15 16.23 16.31 16.38	19.78 19.87 19.96 20.06	15.70 15.76 15.84 15.90
0.9 8 50	8.64 8.71 8.70	10.05	8.52 8.58 8.66 8.73	o.9809 8 7	12.38	15.30	12.14 12.22 12.30 12.37	0.9759 8 7	16.46	20.24	1 2 2
7 6 5 4 3 2	9.00	11.08 11.17 11.26 11.35 11.44	8.80 8.87 8.93 9.00	6 5 4 3 2		15.68 15.77 15.86 15.96 16.05	12.44 12.51 12.59 12.66	6 5 4 3	16.77 16.85 16.92 17.00	20.61 20.71 20.80 20.89 20.99	16.35 16.43
o.9849	9.29 9.36 9-43	11.52 11.61	9.14 9.22 9.29	0.9799	13.08 13.15 13.23	16.15 16.24	12.74 12.81 12.89	2 1 0 0.9749	17.17 17.25	21.09 21.19 21.29	16.74 16.81 16.89
8 7 6 5 4	9.50 9.57 9.64 9.71 9.79	11.79 11.87 11.96 12.05 12.13	9-35 9-42 9-49 9-56 9-64	8 7 6 5 4	13.31 13.38 13.46 13.54 13.62	16.52	13.03 13.10 13.18 13.26	8 7 6 5 4	17.42 17.50 17.58 17.67	21.39 21.49 21.59 21.69 21.79	16.97 17.05 17.13 17.20 17.29
	9.86 9.93 10.00 10.03	12.22 12.31 12.40 12.49	9-71 9-77 9-84 9-92	3 2 1 0	13.69 13.77 13.85 13.92	16.89 16.98 17.08 17.17	13.40 13.48 13.56 13.63	3 2 1	17.83 17.92 18.00 18.08	21.89 21.99 22.09 22.18	17.37 17.46 17.54 17.61
7 ¹	10.15 10.23 10.31 10.38 10.46	12.58 12.68 12.77 12.87 12.96	9.99 10.06 10.13 10.20 10.28	0.9789 8 7 6	14.00 14.09 14.18 14.27 14.36	17.26 17.37 17.48 17.59	13.71 13.79 13.88 13.96 14.04	0.9739 8 7 6	18.15 18.23 18.31 18.38 18.46	22.27 22.36 22.46 22.55 22.64	17.68 17.76 17.82 17.90
4 3 2	10.54 10.62 10.69 10.77	13.05 13.15 13.24 13.34	10.36 10.44 10.51 10.59	4 3 2 1	14.45 14.55 14.64 14.73	17.81 17.92 18.03 18.14	14.13 14.23 14.32 14.39	5 4 3 2 1	18.54 18.62 18.69 18.77	22.73 22.82 22.92 23.01	18.05 18.13 18.19 18.27
<u> </u>	10.85	13-43	10.67	0	14.82	18.25	14.48	٥	18.85	21.10	18.34

	Abe	olute Ak	ohol.		Abe	olute Alc	cohol		Abso	lute Alo	ohoL
Spec. Grav. at 15.6° C.	Per Cent by Weight	Per Cent by Vol- ume.	Grams per 100 cc.	Spec. Grav. at 15.6° C.	Per Cent by Weight	Per Cent by Vol- ume.	Grams per 100 cc.	Spec. Grav. at :5.6° C.	Per Cent by Weight	Per Cent by Voi- ums.	Grams per 100 cc.
0-9729	18.92		18.41	0.9679	22.92		22.18	c 9	26.60		25-61
7	19.00		18.48 18.56		23.00 23.88		22.26	7	26.73	32.42	
် စ်	19.17	23.48	18.65	βó	23.15	28,22	22.40	ó			
5	19.25	23.58	18.73	s	23.23		22.47) s	26.87	32.58	
4	19-33	23.68	18.80	4	23.31		22.54	4	26.93		25.91
3	19.42	23.78	18.88	3	23.38	28.50] 3		32-73	25.98
7	19.50 19.58		18.95	3	23.46 23.54		22.69 22.76	1 3	_,_,	32.81	
ō		24.08	19.12	۰	100		22.83	٥		32.98	26.17
0.9719		24.18	19.19	0.9669		28.86		0.9619		33.06	
9	19.83		19.27	8	23-77 23.85	28.95			27.36		
6	20.00		19.30	6		-		∥ 6			
5	20.08			5	24.00	_	23.111	5	27-57		
4	20.17	24.68	19.59	4	24.08	29.31		4	27.64	33-48	26.57
3	20.25			3			23-33	3	27-71	33-50	26.64
2	20.33		19.74		24-23		23.40	2	1 -1.88		
	20.42		19.83		-4.0		23.48 23-55	6		33-73 33-81	26.78 26.84
0.9709			19.98	0.9659		29.76	23.62	0.9609			26.90
	20.67		-		24.54				28.06	00 / 1	26.96
6	20.75		20.14	6	24.62		23.77 23.84	6	I –		27.01
5	20.92		20.30	5	24.77		23.91	5	28.25	34.18	
4	21.00	25.67	20.33	16	24.85	30.22	23-99	4	28.31	34.25	27.18
3			20.46	3				3	28.37		
2				2		30.40	24.12	2	1 ^	, -	
ô	_			0			24.19 24.26	0		34-47 34-54	
0.969 9	21.38	26.13		0.9649			24.32	0.9599	28.62		27-47
0	21.40		20.81	∥ ౖ*	25.10	30.73 30.82	24-39	8	28.69 28.75	34.69 34.76	27-53
7	21.54		20.96	6	25.36 25.43		,	7		34-70	27.59 27.64
	21.69		21.03	5	3 .0			5	1 - 0.	34.90	
4	21.77	26.58	21.11	∥ 4	25-57	31.07	24.66	4	28.94	34-97	27.76
3				3			24.72	3	29.00		27.82
3	21.92		_	2 1		31.23		3			
•				هٔ			24.93	٥	- × ×		28.00
o. 9689	22.15			0.9639	25.93			0.9589			
8	22.23			4	26.00		25.06	8	, , ,		
6	22.31 22.38	27.22 27.31	21.61	6	26.13	31.65	25.12 25.18	6	29.40 29-47		
5	22.46	27.40	21.76	5	26.20		25.23	5			28.30
4	22.54		21.83	l ă	26.27	31.88	25.30	∥ 4	29,60	35 - 74	28.36
i i	22,62	27-59	21.90	3	26.33	31.96	25.36	3	29.67	35.81	28.43
2	22.69		21.96	2	26.40			2			
1	22.77	27.77 27.86		0			25.49	0	ı		
0	22.05	47.00	22.12	ll Y	26.53	24.42	25-55	II .	1-2.01	30.04	20.01

0	Abs	olute Alc	ohol.	8	Abs	olute Ak	ohol.	Absolute Alcoho			ohoL
Spec. Grav. at 15.6° C.	Per Cent by Weight	Per Cent by Vol- ume.	Grams per 100 cc.	Spec. Grav. at 15.6° C.	Per Cent by Weight	Per Cent by Vol- ume.	Grams per 100 cc.	Spec. Grav. at 15.6° C.	Per Cent by Weight	Per Cent by Vol- ume.	Grams per 100 cc.
0.9579	29.93	36.12	28.67	0.9529	32.94	39.54	31.38	0.9479	35-55	42.45	33-70
8	30.00	36.20	28.73	8	33.00	39.61	31.43	8	35.60	42.51	33-75
7	30.06		28.78	7	33.06	39.68	31.48	7	35.65	42.50	33-79
6	30.11	36.32 36.39	28.82 28.88	6	33.12	39 - 74	31.53	6	35.70	42.62	33.83 33.88
5 4	30.17		28.92	5	33.18 33.24	39.81 39.87	31.59 31.63	5 4	35 - 75 35 - 80	42.67	33.92
3	30.28		28.98	∥ 3 ³	33.29	39.94	31.69	3	35.85	42.78	33.97
2	30.33		29.03	2	33-35	40.01	31.74	2	35.90	42.84	34.01
1	30.39		29.08	1	33.41	40.07	31.80	1	35-95	42.89	34.05
0	30.44		29.13	•	33-47	40.14	31.86	•	36.∞	42.95	34.09
5.956 9	30.50	36.76	29.18	0.9519	-33-53	40.20	31.91	0.9469	36.06		34-14
8	30.56	36.83	29.23	8	33.59	40.27	31.96	8	36.11	43.07	34.09
7 6	30.61		29.27	7 6	33.65	40.34	32.01	7	36.17		34.24
5	30.67		29.33 29.38	5	33-71 33-76	40.40	32.07	5	36.22 36.28	43-19 43.26	34.28 34.34
4	30.78	37.08	29.43	4	33.82	40.53	32.17	4	36.33	43.32	34.38
3	30.83	37.14	29.48	3		40.60	32.22	3	36.39	43.38	34-44
2	30.89	37.20	29.53	∥ ž		40.67	32.27	2	36.44	43-44	34.48
I			29.58	1	34.00	40.74	32.32	1	36.50	43.50	34-54
0	31.00	37-34	29.63	٥	34.05	40.79	32-37	•	36.56	43-56	34.58
0. 9559		37.41	29.69	0.9509	34.10		32.41	0.9459	36.61	43.63	34.63
8	31.12		29.74	8	34.14		32.45	8	36.67	43.69	34.69
6	31.19		29.81	7 6	34.19 34.24		32.49	7 6	36.72 36.78		34 · 73
5	31.31		20.01	5			32.59	5	36.83	43.87	34.83
4				4			32.63	4		43.93	34.88
3			30.03	3			32.67	3			34.92
2		37.90	30.09	2	34-43	41.21	32.71	2	37.00	44.06	34.96
I	10 5	37.97	30.14	1				I	1 37	44.12	35.02
0	31.62	1	١	٥	34-52	41.32	32-79	٥	37-11	44.18	35.07
0.9∄4 9			30.26	0.9499		41.37	32.84	0.9449		44.24	
7	31.75		30.31	H	34.62 34.67		32.88	"	37-22 37-28		
6			30.42	7 6	34.71		32.96	6	37-33		35.26
5	, ,	38.40	30.48	5		41.58	33.00	ll 5	37-39		35-31
4	32.00	38.47	30.53	4	34.81	41.63	33.04	4			35-35
3		38.53	30.59	3			33.09	3			35-41
2	32.12		30.64	2	37-7		33.13	2		44.67	35.40
I	, ,	38.68	30.71	1	1 0. 7		33.17	I o	1 37 -	44-73	35.5
٥	32.25		30-77	l	35.00	41.84	33.21	∥ '	37.67		35.50
0.9 539 8			30.81	0.9489	35.05		32.26	0.9439 8		44.86	35.60
7	32.37		30.87	7	35.10	41.95	33.30	7	37.78 37.83	44.92	35 - 65 35 - 79
6	32.50		30.99	6	35.20		33-34	6	37.89	45.04	35 - 75
5	32.56		31.05	5	35-25	42.12	33.43	5	37.49	45.10	35.80
4	32.62		31.10	4	35.30	42.17	33.48	4	38.00	45.16	35.85
3	32.69	39-25	31.15	3		42.23	33-53	3	38.06	45.22	35.90
2	32.75		31.20	2	35.40	42.29	33-57	2	38.11	45.28	35-95
1	32.81	39.40	31.26	I	35-45	42-34	33.61	0	38.17	45-34	36.∞
0	32.87	39-47	31.32	°	35.50	42.40	33.65	l) °	38.22	45-41	36.04

_	Abs	olute Ak	ohol.		Abs	olute Ale	ohoL		Abso	lute Alc	ohoL
Spec. Grav. at 15.6° C.	Per Cent by Weight	Per Cent by Vol- ume.	Grams per 100 cc.	Spec. Grav. at 15.6° C.	Per Cent by Weight	Per Cent by Vol- ume.	Grams per 100 cc.	Spec. Grav. at 15.6° C.	Per Cent by Weight	Per Cent by Vol- ume.	Gram per
D.94 29	38.28	45-47	36.08	0.9379	40.85	48.26	38.31	0.9329	43.29	50.87	40.3
8	38.33	45-53	36.13	8	40.90		38.35	8	43-33	50.92	40.4
7	38.39	45-59	36.18	7	40.95	48.37	38.39	7	43-39	50.97	40.4
6	38.44		36.23	6	41.00	48.43	38.44	6	43.43	51.02	40.5
5	38.50	45-71	36.28	5	41.05	48.48	38.48	5	43.48	51.07	40.5
4	38.56		36.33	4	41.10		38.52	4	43.52	51.12	40.5
3	38.61 38.67	45.83 45.89	36.38	3	41.15	48.59 48.64	38.58 38.62	3	43.57	51.17	40.6
1	38.72	,	36.43 36.48	2 1	41.20	48.70	38.66	2 I	43.62		40.6
ò	38.78		36.53	6	41.30		38.70	o	43.67 43.71	51.27 51.32	40.7
0.9 419	38.83		36.57	0.9369	41.35	48.80	38.74	0.9319	43.76	51.38	40.7
8	38.89		36.62	8	41.40		38.78	8	43.81	51.43	40.8
7	38.94		36.67	7	41.45	48.91	38.82	7	43.86		40.8
6	39.00	46.26	36.72 36.76	11	41.50	48.97	38.87	6	43.90		40.8
5	39.05	46.32	36.80	5	41.55	49.02	38.91	5	43-95	51.58	40.9
4		46.42	36.85	3	41.65	49.13	38.95 38.99	4	44.00	51.63 51.68	40.9
3 2	39.20		36.80	3	41.70		39.04	3 2	44.00	51.72	41.0
1	39.25	46.53	36.94	1	41.75	49.23	39.08	ī	44.14	51.77	41.0
0			36.98	0	41.80		39.13	0	44.18	51.82	41 1
o 940 9	39-35		37.02	0.9359	41.85		39.17	0.9309	44-23	51.87	41.1
	39.40		37.07	8	41.90		39.21	8	44-27	51.91	41.2
7 6	39-45	46.75 46.80	37.11	7	41.95	49-45	39.25	7 6	44-32	51.96	41.2
	39-50	46.86	37.15	5	42.00	49 50 49 55	39.30 39.34		44.30	52.01 52.06	41.2
5 4	39·55 39.60	46.91	37-23	4	42.10		39.38	5 4	44.41 44.46	52.10	41.3
3	39.65	46.97	37-27	3	42.14		39.42	3	44.50		41.4
2			37-32	2	42.19		39.46	2	44-55	52.20	41.4
1	39-75		37.36	1	42.24	49.76	39.50	1	44.59	52.25	41.4
0	39.80	47-13	37-41	∥ °	42.29	49.81	39-54	٥	44.64	52.29	41.5
9.9 399	39.85		37-45	0.9349	42.33 42.38	49.86 49.91	39.58 39.62	0.9299 8	44.68	52.34	41.5 41.5
7	39.90		37-49 37-53	7	42.43		39.66	7	44-73	52.39 52.44	41.6
6			37.58	∥ 6			39.70	6	44.82	52.48	41.0
5	, -	1	37.62	5	42.52		39-74	5	44.86		41.7
4			37.67		42.57	50.11	39.78	4	44.91	52.58	41.
3		47.51	37-71	3	42.62		39.82	3	44.96	52.63	41.
2	40.20		37-75	2		50.21	39.86	2	45.00	, ,	41.8
1	, , ,		37.80	1	42.71		39.90	1	45.05	52.72	41.
0	40.30	47.67	37.84	°	42.76	50.31	39-94	°	45.09	52-77	41.8
9.93 89 8	40.35		37.88 37.92	0.9339	42.81 42.86	50-37 50-42	39.98 40.02	0.9289	45-14 45.18		41.9
7	40.45	47.83	37.96	7	42.90		40.06	7	45.23	52.91	42.0
é			38.00	6	42.95	50.52	40.10	6	45.27	52.96	42.0
5	40.55	47-94	38.05	5	43.00		40.14	5	45-32	53.01	42.0
4	40.60	47.99	38.09	4	43.05	50.62	40.18	4	45.36	53.06	42.1
3	40.65	48.05	38.13	3	43.10		40.22	3	45.41	53.10	42.
2	40.70		38.18	2	43.13	50.72	40.26	2	45.46		42.1
I	40.75	48.16	38.22	1	43.19		40.30	1	45-50		42.2
0	40.80	48.21	38.27	•	43-24	50.82	40.34	•	45.55	53.24	42.2

Sec.	Absolute	Alcohol.	Se	Absolute	Alcohol	Spec.	Absolute	Alcohol.
Spec. Grav. at 15.6° C.	Per Cent	Per	Spec. Grav. at 15.6° C.	Per Cent	Per	Grav. at 15.6° C.	Per Cent	Per Cent
15.0° C.	by Weight.	Cent by Vol- ume.	15.0° C.	by Weight.	Cent by Vol- ume.	15.0 %	by Weight.	by Vol- ume.
0.927 9 8	45-59	53-29	0.9229	47-86	55-65	0.9179	50.13	57.97
	45.64 45.68	53-34	8	47.91	55.69	11	50.17	58.01 58.06
7	45.73	53·39 53·43	7 6	47.96 48.∞	55·74 55·79	7 6	50.22 50.26	58.10
5	45.77	53.48	5	48.05	55.83	5	50.30	58.14
4	45.82	53-53	3	48.00	55.88	4	50.35	58.19
3	45.86	53.58	3	48.14	55-93	∥ <u>š</u>	50.39	58.23
2	45.91	53.62	2	48.18	55-97	2	50.43	58.28
1	45.96	53.67	1	48.23	56.02	1	50.48	58.32
•	46.00	53-72	•	48.27	56.07	۰	50.52	58.36
0.9269	46.05	53-77	0.9219	48.32	56.11	0.9169	50-57	58.41
8	46.09	53.81	8	48.36	56.16	8	50.61	58.45
7 6	46.14 46.18	53.86	7 6	48.41	56.21	7	50.65	58.50
	46.23	53.91	11	48.46	56.25	11	50.70	58-54
5 4	46.27	53-95 54.00	5 4	48.50	56.30	5 4	50.74 50.78	58.58 58.63
3	46.32	54.05	3	48.55 48.59	56.40	3	50.83	58.67
2	46.36	54.10	2	48.64	56.44	2	50.87	58.72
1	46.41	54.14	ī	48.68	56.40	ī	50.91	58.76
•	46.46	54-19	-	48.73	56.54	0	50.96	58.80
0.9259	46.50	54-24	0.9209	48.77	56.58	0.9159	51.00	58.85
8	46.55	54-29	8	48.82	56.63	8	51.04	58.89
7 6	46.59	54-33	7	48.86	56.68	7	51.08	58.93
	46.64	54-38	6	48.91	56,72	6	51.13	58.97
5	46.68	54-43	5	48.96	56.77	5	51.17	59.01
4	46.73	54-47	4	49.00	56.82	4	51.21	59.05
3	46.82	54-52	3	49.04	56.8 6	3	51.25	59.09
1	46.86	54.57 54.62	2	49.08	56.90	2	51.29	59.14
ō	46.91	54.66		49.16	56.94 56.98	0	51-33 51.38	59.18 59.22
0.9249	46.96	54-71	0.9199	49.20	57.02	0.9149	51.42	59.26
8	47.00	54.76	Proof 8	49.24	57.06	8	51.46	59.30
7 6	47.05	54.80	7	49.29	57.10	7 6	51.50	59-34
	47.00	54.85	6	49-34	57-15	11	51.54	59.39
5	47.14	54.90	5	49.39	57.20	5	51.58	59-43
4	47.18	54.95	4	49.44	57-25	4	51.63	59-47
3	47.27	54-99 55-04	3	49-49	57-30	3	51.67	59.51
ī	47.32	55.09	2	49.54	57-35	2	51.71	59.55
ō	47.36	55.13	6	49.59 49.64	57.40	0	51.75 51.79	59.59 59.63
0.9239	47-41	55.18	0.9189	49.68	57-49	0.9139	51.83	59.68
8	47.46	55-23	8	49-73	57.54	373.38	51.88	59-72
7 6	47-50	55-27	7	49.77	57-59	7	51.92	59.76
	47.55	55-32	6	49.82	57.64	6	51.96	59.80
5	47.59	55-37	5	49.86	57.69	5	52.00	59.84
4	47.64	55-41	4	49.91	57-74	4	52.05	59.89
3	47.68	55.46	3	49-95	57-79	3	52.09	59.93
2 I	47.73	55-51	2	50.00	57.84	2	52.14	59.98
0	47-77 47-82	55.55	1	50.04	57.88	1	52.18	60.02
-	4,.02	55.60	0	50.09	58.92	0	52.23	60.07

ALCOHOLIC BEVERAGES.

_	Absolute	Alcohol.		Absolute	Alcohol		Absolute	Alcohol.
Spec. Grav.	Per	Per	Spec. Grav.	Per	Per	Spec. Grav.	Per	Per
15.6° C.	Cent	Cent by Vol-	15.6° C.	Cent	Cent	at 15.6° C.	Cent	Cent
13.0 .	by Weight.	by Vol- ume.	15.0 .	by Weight.	Cent by Vol- ume.	13.0 .	by Weight.	by Vol- ume.
0.0120	52.27	60.12	0.9079	54-52	62.36	0.9029	56.82	64.63
	52.32	60.16	8	54.57	62.41	8	56.86	64.67
7	52.36	60.21	7	54.62	62.45	7	56.91	64.71
7 6	52.41	60.25	6	54.67	62.50	6	56.95	64.76
5	52-45	60.30	5	54-71	62.55	5	57.00	64.80
4	52.50	60.34	4	54.76	62.60	4	57.04	64.85
3	52-55	60.39	3	54.81	62.65	3	57.08	64.89
2	52-59	60.44	2	54.86	62.69	2	57-13	64.93
I	52.64	60.47	I	54.90	62.74	I	57.17	64.97
0	52.68	60.52	۰	54-95	62.79	٥	57.21	65.01
0.9119	52.73	60.56	0.9069	55.00	62.84	0.9019	57-25	65.05
8	52-77	60.61	8	55.05	62.88	8	57.29	65.09
7 6	52.82	60.65	7	55.09	62.93	7	57-33	65.13
	52.86	60.70	6	55-14	62.97	6	57-38	65.17
5	52.91	60.74	5	55.18	63.02	5	57.42	65.21
4	52.95	60.79	4	55-23	63.06	4	57.46	65.25
3	53.00	60.85	3	55-27	63.11	3	57-50	65.29
2	53.04	60.89	2	55-32	63.15	2	57.54	65.33
0	53.09 53.13	60.93	1	55.30 55.41	63.20		57.58 57.63	65.37 65.41
	1					1		
0.910 9	53-17	61.02	0.9059	55-45	63.28	0.9009	57.67	65.45
	53.22 53.26	61.10		55-50	63.33	11	57-71 57-75	65.53
7	53.30	61.15	7 6	55.55	63.37	7 6	57.79	65.57
5	53-35	61.10	5	55.59 55.64	63.46	5	57.83	65.61
4	53-39	61.23	4	55.68	63.51	4	57.88	65.65
3	53.43	61.28	3	55.73	63.55	3	57.92	65.69
2	53.48	61.32	3	55-77	63.60	2	57.96	65.73
1	53-52	61.36	1 1	55.82	63.64	1	58.00	65.77
0	53-57	61.40	•	55.86	63.69	•	58.05	65.81
0.9 099	53.61	61.45	0.9049	55.91	63.73	0.8999	58.09	65.85
8	53.65	61.49	8	55-95	63.78	8	58.14	65.90
7	53-70	61.53	7 6	56.00	63.82	7	58.18	65.94
6	53-74	61.58	11	56.05	63.87	6	58.23	65.99
5	53.78	61.62	5	56.09	63.91	5	58.27	66.03
4	53.83	61.66	4	56.14	63.96	4	58.32 58.36	66.12
3	53.87	61.71	3 2	56.18	64.00	3	58.41	66.16
2 I	53.91 53.96	61.75	1 1	56.23	64.05 64.09	ī	58.45	66.21
0	54.00	61.84		56.32	64.14		58.50	66.25
0.908 9	54.05	61.88	0 0030	56.36	64.18	0.8989	58.55	66.29
8	54.10	61.03	0.9039	56.41	64.22	0.0909	58.59	66.34
7	54.14	61.98	7	56.45	64.27	7	58.64	66.38
6	54.19	62.03	6	56.30	64.31	6	58.68	66.43
5	54.24	62.07	5	56.55	64.36	5	58.73	66.47
4	54.29	62.12	4	56.59	64.40	4	58.77	66.51
3	54-33	62.17	3	56.64	64.45	3	58.82	66.56
2	54.38	62.22	2	56.68	64.49	2	58.86	66.60
I	54-43	62.26	I	56.73	64.54	1	58.91	66.65
0	54.48	62.31	ه اا	56.77	64.58	li o	58.95	66.60

S	Absolute	Alcohol.		Absolute	AlcohoL	S	Absolute	Alcohol.
Spec. Grav. at 15.6° C.	Per Cent by Weight.	Per Cent by Vol- ume.	Spec. Grav. at 15.6° C.	Per Cent by Weight.	Per Cent by Vol- ume.	Spec. Grav. at 15.6° C.	Per Cent by Weight.	Per Cent by Vol- ume.
0.8979	59.00	66.74	0.8929	61.13	68.76	0.8879	63.30	70.81
8	59.04	66.78	8	61.17	68.80	8	63.35	70.85
7 6	59.09	66.82	7 6	61.21	68.83 68.87	7 6	63.39	70.89
5	59.13 59.17	66.90	5	61.25 61.29	68.91	5	63.43 63.48	70.93
4	59.22	66.94	4	61.33	68.95	4	63.52	71.01
3	59.26	66.99	š	61.38	68.99	3	63.57	71.05
2	59.30	67.03	2	61.42	69.03	2	63.61	71.09
1	59-35	67.07	ɪ	61.46	69.07	1	63.65	71.13
0	59-39	67.11	•	61.50	69.11	٥	63.70	71.17
0.8969	59.43	67.15	0.8919	61.54	69.15	0.8869	63.74	71.22
8	59.48	67.19	8	61.58	69.19	8	63.78	71.26
7	59.52	67.24	7 6	61.63 61.67	69.22	7 6	63.83	71.30
5	59·57 5 9 ·61	67.32	5	61.71	69.30	5	63.87 63.91	71.34
4	59.65	67.36	4	61.75	69.34	4	63.96	71.42
3	59.70	67.40	3	61.79	69.38	3	64.00	71.46
2	59-74	67.44	2	61.83	69.42	2	64.04	71.50
I	59.78	67.49	l r	61.88	69.46	1	64.09	71.54
0	59.83	67.53	•	61.92	69.50	•	64.13	71.58
0.8959	59.87	67.57	0.8909	61.96	69.54	o.8859 8	64.17	71.62
8	59.91	67.61	8	62.00	69.58		64.22	71.66
7 6	59.96 60.00	67.65	7 6	62.05	69.62	7 6	64.26	71.70
5	60.04	67.73	5	62.09 62.14	69.71	5	64.30 64.35	71.74
4	60.08	67.77	4	62.18	69.75	4	64.39	71.82
3	60.13	67.81	3	62.23	69.79	3	64.43	71.86
2	60.17	67.85	2	62.27	69.84	2	64.48	71.90
1	60.21	67.89	I	62.32	69.88	1	64.52	71.94
0	60.26	67.93	•	62.36	69.92	•	64.57	71.98
6.8 949	60.29	67.97	0.8899	62.41	69.96	0.8849	64.61	72.02
8	60.33	68.01 68.05	8	62.45	70.01	8	64.65	72.00
7 6	60.38 60.42	68.00	7 6	62.50 62.55	70.05	7 6	64.70 64.74	72.10
5	60.46	68.13	5	62.59	70.14	5	64.78	72.18
4	60.50	68.17	4	62.64	70.18	4	64.83	72.22
3	60.54	68.21	3	62.68	70.22	3	64.87	72.20
2	60.58	68.25	2	62.73	70.27] 2	64.91	72.30
I	60.63	68 29	I	62.77	70.31	I	64.96	72.34
0	60.67	68.33	•	62.82	70-35	•	65.00	72.38
0.8 939	60.71	68.36	0.8889	62.86	70.40	0.8839	65.04	72.42
8	60.76	68.40	8	62.91	70.44	8	65.08	72.40
7 6	60.79 60.83	68.44 68.48	7 6	62.95 63.00	70.48	7 6	65.13	72.50
5	60.88	68.52	5	63.04	70.52	5	65.21	72.58
4	60.92	68.56	3 4	63.09	70.61	3 4	65.25	72.61
3	60.96	68.60	3	63.13	70.65	3	65.29	72.65
2	61.00	68.64	2	63.17	70.69	3	65.33	72.69
1	61.04	68.68	1	63.22	70.73	r	65.38	72.73
0	61.08	68.72	0	63.26	70-77	0	65.42	72.77

ALCOHOLIC BEVERAGES.

						,	· · · · · · · · · · · · · · · · · · ·	
Spec.	Absolute	Alcohol.	Spec.	Absolute	Alcohol.	Spec.	Absolute	Alcohol.
Grav.	Per	Per	Grav.	Per	Per	Grav.	Per	Per
15.6° C.	Cent	Cent by Vol-	25.6° C.	Cent	Cent by Vol-	15.6° C.	Cent by	Cent by Vol-
	Weight.	ume.		Weight.	ume.		Weight.	ume.
0.8829	65.46	72.80	0.8779	67.58	74-74	0.8729	69.67	76.6r
8	65.50	72.84	8	67.63	74-78	8	69.71	76.65
7 6	65.54	72.88	7 6	67.67 67.71	74.82 74.86	7 6	69.75 69.79	76.68
5	65.58 65.63	72.96	5	67.75	74.89	5	69.83	76.72 76.76
4	65.67	72.99	4	67.79	74-93	4	69.88	76.80
3	65.71	73.03	3	67.79 67.83	74-97	3	69.92	76.83
2	65.75	73-07	2	67.88	75.01	2	69.96	76.87
I	65.79	73.11	I	67.92	75.04	I	70.00	76.91
•	65.83	73-15	•	67.96	75.08	٥	70.04	76.94
o.8819	65.88	73.19	0.8769	68.00	75.12	0.8719	70.08	76.98
8	65.92	73.22	8	68.04 68.08	75.16	8	70.12	77.01
7 6	65.96 66.00	73.26 73.30	6	68.13	75.19 75.23	7.	70.16 70.20	77.05 77.08
5	66.04	73-34	5	68.17	75.27	5	70.24	77.12
4	66.09	73.38	4	68.21	75-30	4	70.28	77-15
3	66.13	73.42	3	68.25	75-34	3	70.32	77.19
2	66.17	73.46	2	68.29	75 - 38	2	70.36	77.22
I	66.22 66.26	73-50	I	68.33 68.38	75-42	I	70.40	77-25
٥	00.20	73-54	•		75-45	•	70.44	77-29
0.880 0	66.30	73.57	0.8759	68.42 68.46	75-49	0.8709	70.48	77-32
-	66.35 66.39	73.61 73.65	7	68.50	75-53	9	70.52 70.56	77-36
7 6	66.43	73.69	7 6	68.54	75-57 75-60	7 6	70.60	77-39 77-43
5	66.48	73-73	5	68.58	75.64	5	70.64	77.46
4	66.52	73-77	4	68.63	75.68	4	70.68	77-50
3	66.57	73.81	3	68.67	75-72	3	70.72	77.53
2	66.61	73.85	2	68.71	75-75	2 1	70.76	77-57
0	66.65 66.70	73-89 73-93	,	68.75 68.79	75.79		70.80 70.84	77.60
. 0				68.83	1	06		
0. 8799 8	66.74 66.78	73-97 74-01	0.8749	68.88	75.87 75.90	o.8699 8	70.88	77.67
7	66.83	74.05	1	68.92	75-94		70.96	77.74
6	66.87	74.09	7 6	68.96	75.98	7 6	71.00	77.78
5	66.91	74-13	5	69.00	76.01	5	71.04	77.82
4	66.96	74-17	4	69.04	76.05	4	71.08	77.85
3	67.00	74.22	3	69.08	76.09	3	71.13	77.89
2 I	67.04 67.08	74-25	2 1	69.13	76.13 76.16	2 1	71.17	77-93
•	67.13	74-29		69.21	76.20		71.25	78.00
0.8789	6			60.00	-6	0.8680	' '	-0
8	67.17 67.21	74-37	0.8739	69.25	76.24 76.27	8	71.29	78.04
_	67.25	74-44	7	69.33	76.31	7	71.38	78.11
7 6	67.29	74.48	6	69.38	76.35	6	71.42	78.14
5	67.33	74-52	5	69.42	76.39	5	71.46	78.18
4	67.38	74-55	4	69.46	76.42	4	71.50	78.22
3	67.42	74.59	3 2	69.50	76.46 76.50	3	71.54	78.25
1	67.50	74.67	1	69.54	76.53	1 1	71.50	78.33
0	67.54	74.70	0	69.63	76.57	0	71.67	78.36
		1	1	1	1	<u> </u>		<u> </u>

Spec.	Absolute	Alcohol.	Spec.	Absolute	Alcohol.	Spec.	Absolute	Alcohol.
Grav.	Per	Per	Grav.	Per	Per	Grav.	Per	Per
at 15.6° C.	Cent	Cent by Vol-	at 15.6° C.	Cent	Cent by Vol-	15.6° C.	Cent	Cent
15.0 C.	by	by Vol-	-3.5	by	by Vol-	-3.0 -	by	by Vol-
	Weight.	ume.		Weight.	ume.		Weight.	ume.
.8679	71.71	78.40	0.8629	73.83	80.26	0.8579	76.08	82.23
8	71.75	78.44	8	73.88	80.30	8	76.13	82.26
7	71-79	78.47	7	73.92	80.33	7	76.17	82.30
6	71.83	78.51	6	73.96	80.37	6	76.21	82.33
5	71.88	78.55	5	74.00	80.40	5	76.25	82.37
4	71.92	78.58	4	74-05	80.44	4	76.29	82.40
3	71.96	78.62	3	74.09	80.48	3	76.33	82.44
2	72.00	78.66	2	74-14	80.52	2	76.38	82.47
I	72.04	78.70	i I	74.18	80.56	l I	76.42	82.51
0	72.09	78.73	•	74.23	80.60	°	76.46	82.54
.8660	72.13	78.77	0.8619	74.27	80.64	0.8569	76.50	82.58
8	72.17	78.8r	8	74 - 32	80.68	8	76.54	82.61
7	72.22	78.85	7	74.36	80.72	7	76.58	82.65
7	72.26	78.89	6	74.41	80.76	7 6	76.63	82.69
5	72.30	78.93	5	74 - 45	80.80	5	76.67	82.72
4	72.35	78.96	4	74.50	80.84	4	76.71	82.76
3	72.39	79.00	3	74-55	80.88	3	76.75	82.79
2	72-43	79.04	2	74-59	80.92	ž	76.79	82.83
1	72.48	79.08	1	74.64	80.96	ı	76.83	82.86
0	72.52	79.12	. •	74.68	81.00	0	76.88	82.90
.8659	72-57	79.16	0.8600	74-73	81.04	0.8559	76.92	82.93
8	72.61	79.19	8	74-77	81.08	°8	76.96	82.97
7	72.65	79.23	7	74.82	81.12	7	77.∞	83.00
6	72.70	79.27	7 6	74.86	81.16	7 6	77.04	83.04
5	72-74	79.31	5	74-91	81.20	5	77.08	83.07
4	72.78	79.35	4	74-95	81.24	4	77-13	83.11
3	72.83	79-39	3	75.00	81.28	3	77.17	83.14
2	72.87	79.42	2	75.05	81.32	3	77.21	83.18
I	72.01	79.46	1	75-09	81.36	r	77-25	83.21
0	72.96	79-50	0	75-14	81.40	0	77-29	83.25
.8649	73.00	79-54	0.8599	75.18	81.44	0.8549	77-33	83.28
8	73.04	79.57	8	75-23	81.48	8	77.38	83.32
7	73.08	79.61	7	75-27	81.52	7	77-42	83.36
7 6	73.13	79.65	6	75-33	81.56	7 6	77.46	83.39
5	73-17	79.68	5	75.36	81.Ğo	5	77.50	83.43
4	73.21	79-72	4	75-41	81.64	4	77-54	83.46
3	73.25	79-75	3	75-45	81.68	3	77.58	83.50
2	73-29	79.79	2	75.50	81.72		77.63	83.53
I	73-33	79.83	1 1	75-55	81.76	1	77.67	83.57
0	73.38	79.86	0	75-59	81.80	0	77.71	83.60
.8639	73-42	79.90	0.8589	75.64	81.84	0.8539	77-75	83.64
8	73.46	79-94	Š	75.68	81.88	8	77 - 79	83.67
	73.50	79-97	7	75.73	81.92	7	77.83	83.7
7	73-54	80.01	7 6	75-77	81.96	7 6	77.88	83.74
5	73.58	80.04	5	75.82	82.00	5	77-92	83.78
4	73.63	80.o8	4	75.86	82.04	4	77.96	83.81
3	73.67	80.12	3	75.91	82.08	3	78.∞	83.89
2	73-71	80.15	2	75-95	82.12	2	78.04	83.88
1	73-75	80.19	1	76.00	82.16	r	78. 0 8	83.91
	73.79	80.22	ا ا	76.04	82.10	ا ہ	78.12	83.94

	Absolute	Alcohol.		Absolute	Alcohol.		Absolute	Alcohol.
Spec. Grav. at 15.6° C.	Per Cent	Per Cent. by Vol-	Spec. Grav. at 15.6° C.	Per Cent	Per Cent by Vol-	Spec. Grav. at 15.6° C.	Per Cent	Per Cent
15.0 0.	by Weight.	by Vol- ume.	13.0 0.	by Weight.	by Vol- ume.		by Weight.	by Vol- ume.
0.8529	78.16	83.98	0.8479	80.17	85.63	0.8429	82.19	87.27
-	78.20 78.24	84.01	II	80.21 80.25	85.66	- 1	82.23 82.27	87.30
7	78.28	84.08	7 6	80.20	85.70 85.73	7 6	82.31	87.34 87.37
5	78.32	84.11	5	80.33	85.77	5	82.35	87.40
4	78.36	84.14	4	80.38	85.77 85.80	4	82.38	87.43
3	78.40	84.18	3	80.42	85.84	3	82.42	87.46
2	78.44	84.21	2	80.46	85.87	2	82.46	87-49
1	78.48	84.24	1	80.50	85.90	I	82.50	87.52
0	78.52	84.27	•	80.54	85.94	0	82.54	87-55
0.8519	78.56	84.31	0.8469	80.58	85.97	0.8419	82.58	87.58
8	78.60	84-34	8	80.63	86.or	8	82.62	87.6r
7	78.64	84.37	7	80.67	86.04	7 6	82.65	87.64
	78.68	84.41	6	80.71	86.08 86.11		82.69	87.67
5	78.72	84.44	5	80.75 80.79	86.15	5	82.73 82.77	87.70
4	78.76 78.80	84.47 84.51	4	80.79	86.18	4	82.81	87.76
3 2	78.84	84.54	3 2	80.88	86.22	3 2	82.85	87.79
ī	78.88	84.57	ī	80.02	86.25	1	82.88	87.82
0	78.92	84.60	0	80.96	86.28	0	82.92	87.85
0.8509	78.96	84.64	0.8459	81.00	86.32	0,8409	82.96	87.88
8	79.00	84.67	8	81.04	86.35	8	83.00	87.91
7	79.04	84.70	7 6	81.08	86.38	7 6	83.04	87.94
	79.08	84-74		81.12	86.42		83.08	87.97
5	79.12	84.77 84.80	5	81.16 81.20	86.45 86.48	5	83.12 83.15	88.00 88.03
4	79.16 79.20	84.83	4 3	81.24	86.51	4 3	83.19	88.06
3	79.24	84.87	2	81.28	86.54	2	83.23	88.00
ī	79.28	84.90	1	81.32	86.58	1	83.27	88.13
0	79-32	84.93	٥	81.36	86.61	٥	83.31	88.16
0.8499	79.36	84.97	0.8449	81.40	86.64	0.8399	83.35	88.19
8	79.40	85.00	8	81.44	86.67	8	83.38	88.22
7	79-44	85.03	7	81.48	86.71	7 6	83.42	88.25 88.28
	79.48	85.06	6	81.52 81.56	86.74 86.77	5	83.46 83.50	88.31
. 5	79.52 79.56	85.10 85.13	5 4	81.60	86.80	3 4	83.54	88.34
4 3	79.50	85.16	3	81.64	86.83	3	83.58	88.37
2	79.64	85.19	2	81.68	86.87	2	83.62	88.40
ī	79.68	85.23	1	81.72	86.90	1	83.65	88.43
ō	79.72	85.26	٥	81.76	86.93	٥	83.69	88.46
0.8489	79.76	85.29	0.8439	81.80	86.96	0.8389	83.73	88.49
8	79.80	85.33	8	81.84	86.99	8	83.77	88.52
7 6	79.84	85.36	7	81.88	87.03	7 6	83.81	88.5 5 88.5 8
	79.88	85.39	6	81.92	87.06		83.85	
5	79.92	85.42	5	81.96 82.00	87.09 87.12	5 4	83.88 83.92	88.61 88.64
4	79.96 80.00	85.46 85.40	4	82.04	87.15	3	83.96	88.67
3	80.04	85.49 85.53	3 2	82.08	87.18	2	84.00	88.70
1	80.08	85.56	1	82.12	87.21	1	84-04	88.73
0	80.13	85.59	0	82.15	87.24	0	84.08	88.76
			J			l		<u> </u>

	Absolute	Alcohol.		Absolute	Alcohol.		Absolute	Alcohol.
Spec. Grav.			Spec. Grav.	n	Per	Spec. Grav.		
15.6° C.	Per Cent	Per Cent	at 15.6° C.	Per Cent	Cent	15.6° C.	Per Cent	Per Cent
15.0 6.	by	by Vol-	15.0° C.	by	by Vol-	15.6° C.	by	Cent. by Vol-
	Weight.	ume.		Weight.	ume.		Weight.	ume.
.8379	84.12	88.79	0.8329	86.08	90.32	0.8279	88.00	91.78
8	84.16	88.83	8	86.12	90.35	8	88.04	91.81
7 6	84.20	88.86 88.80	7 6	86.15 86.19	90.38	7 6	88.08 88.12	91.84
	84.24	88.92	11	86.23	90.40		88.16	91.87
5	84.28 84.32	88.95	5	86.27	90.43	5	88.20	91.90
4 3	84.36	88.98	4	86.31	90.40	4	88.24	91.93 91.96
3 2	84.40	89.01	3 2	86.35	90.52	3 2	88.28	91.99
1	84.44	80.05	ī	86.38	90.55	ī	88.32	92.02
ō	84.48	89.08	ō	86.42	90.58	ò	88.36	92.05
9.8 369	84.52	89.11	0.8319	86.46	90.61	0.8269	88.40	92.08
ž g	84.56	80.14	8	86.50	90.64	8	88.44	92.12
7	84.60	89.17	7	86.54	90.67	7	88.48	92.15
6	84.64	89.20	6	86.58	90.70	6	88.52	92.18
5	84.68	89.24	5	86.62	90.73	5	88.56	92.21
4	84.72	89.27	4	86.65	90.76	i 4	88.60	92.24
3	84.76	89.30	3	86.69	90.79	3	88.64	92.27
2	84.80	89.33	2	86.73	90.82	2	88.68	92.30
I	84.84	89.36	1	86.77	90.85	' I	88.72	92.33
0	84.88	89.39	•	86.81	90.88	•	88.76	92.36
0.8 359	84.92	89.42	0.8309	86.85	90.90	0.8259	88.8o	92.39
8	84.96	89.46	8	86.88	90.93	8	88.84	92.42
7	85.00	89.49	7 6	86.92	90.96	7	88.88	92.45
6	85.04	89.52		86.96	90.99	6	88.92	92.48
5	85.08	89.55	5	87.00	91.02	5	88.96	92-51
4	85.12	89.58	4	87.04	91.05	4	89.00	92.54
3	85.15	89.61	3	87.08	91.08	3	89.04	92.57
2	85.19	89.64	2	87.12	91.11	2	89.08	92.60
I 0	85.23 85.27	89.67 89.70	1 0	87.15 87.19	91.14	1 0	89.12 89.16	92.6
					' '			'
0.8 349 8	85.31	89.72	0.8299	87.23	91.20	0.8249	89.19	92.68
	85.35	89.75 89.78		87.27	91.23	8	89.23	92.71
7 6	85.38 85.42	80.81	7 6	87.31	91.25	7 6	89.27	92.74
5	85.46	89.84	5	87.35 87.38	91.28	11	89.31	92.77
4	85.50	89.87	3	87.42	91.31	5	89.35 89.38	92.8
3	85.54	89.90	3	87.46	91.34	4	89.42	92.8
2	85.58	89.93	3	87.50	91.40	3 2	89.46	92.80
ī	85.62	89.96	ī	87.54	91.43	1	89.50	92.0
ō	85.65	89.99	0	87.58	91.46	0	89.54	92.9
0.8339	85.69	90.02	0.8280	87.62	91.49	0.8239	89.58	92.9
8	85.73	90.05	8	87.65	91.52	8	89.62	93.00
7 6	85.77	90.08	7 6	87.69	91.55	7	89.65	93.0
6	85.81	90.11	6	87.73	91.57	6	89.69	93.0
5	85.85	90.14	5	87.77	91.60	5	89.73	93.0
4	85.88	90.17	4	87.81	91.63	4	89.77	93.1
3	85.92	90.20	3	87.85	91.66	3	89.81	93.1
2	85.96	90.23	2	87.88	91.69	2	89.85	93.1
I	86.00	90.26	I	87.92	91.72	1	89.88	93.20
0	86.04	90.29	' 0	87.96	91.75	0	89.92	93.23

Spec.	Absolute	Alcohol.	Spec.	Absolute	Alcohol.	Spec.	Absolute	AlcohoL
Grav.	Per	Per	Grav.	Per	Per	Grav.	Per	D
at 15.6° C.	Cent by	Cent by Vol-	15.6° C.	Cent	Cent by Vol-	at 15.6° C.	Cent	Per Cent
23.0	Weight.	by Vol- ume.	13.0 0.	by Weight.	by Vol- ume.	13.0 0.	by Weight.	Cent by Vol- ume.
0.8229	89.96	93.26	0.8179	91.75	94-53	0.8120	93.59	95.84
8	90.00	93.29	8	91.79	94.56	8	93.63	95.87
7 6	90.04	93.31	7	91.82	94-59	7	93.67	95.90
	90.07	93.34	6	91.86	94.61	6	93.70	95.92
5	90.11	93.36	5	91.89	94.64	5	93-74	95.95
4	90.14	93-39	† 4	91.93	94.66	4	93.78	95-97
3	90.18	93.41	3	91.96	94.69	3	93.81	96.00
2 I	90.21	93-44	2	92.00	94.71	2	93.85	96.03
0	90.25 90.29	93-47	I 0	92.04	94 - 74	I	93.89	96.05
	90.29	93-49		92.07	94.76	0	93-92	96.08
0.8219	90.32	93-52	0.8169	92.11	94-79	0.8119	93.96	96.11
8	90.36	93-74	8	92.15	94.82	8	94.00	96.13
7 6	90.39	93-57	7	92.18	94.84	7	94.03	96.16
	90.43	93-59	6	92.22	94.87	6	94.07	96.18
5	90.46	93.62	5	92.26	94.90	5	94.10	96.20
4	90.50	93.64	4	92.30	94-92	4	94-14	96.22
3	90.54 90.57	93.67	3	92.33	94-95	3	94.17	96.25
1	90.57	93.72	2 I	92.37	94.98	2 1	94.21	96.27
0	90.64	93.72		92.41 92.44	95.00 95.03	0	94.24 94.28	96.29
	(0						94.20	90.3-
0. 8200	90.68	93-77	0.8159	92.48	95.06	0.8109	94.31	96.34
- 1	90.71	93.80	8	92.52	95.08	8	94-34	96.36
7	90.75	93.82	7	92.55	95.11	7	94.38	96.39
5	90.79 90.82	93.85	6	92.59	95.13	6	94.41	96.41
4	9c.86	93.90	5	92.63 92.67	95.16	5	94.45	96.43
3	90.89	93.93	4 3	92.70	95.19 95-21	4 3	94.48	96.46
2	90.93	93.95	2	92.74	95-24	2	94 - 52 94 - 55	96.50
I	90.96	93.98	1	92.78	95.27		94-55	96.53
0	91.00	94.00	0	92.81	95.29	0	94.62	96.55
0.8199	91.04	94.03	0.8149	92.85	95-32	0.8000	94.65	96.57
8	91.07	94.05	8	92.89	95-35	8	94.69	96.60
7	91.11	94.08	7	92.92	95 - 37	7	94-73	96.62
6	91.14	94.10	6	92.96	95.40	6	94.76	96.64
5	91.18	94.13	5	93.00	95.42	5	94.80	96.67
4	91.21	94.15	4	93.04	95-45	4	94.83	96.69
3	91.25	94.18	3	93.07	95.48	3	94.86	96.71
2	91.29	94.21	2	93.11	95.50	2	94.90	96.74
I	91.32	94.23	I	93.15	95-53	I	94-93	96.76
0	91.36	94.26		93.18	95-55	•	94-97	96.78
0.8189	91.39	94.28	0.8139	93.22	95.58	0.8089	95.00	96.80
8	91.43	94.31	8	93.26	95.61	8	95.04	96.83
7	91.46	94-33	7	93.30	95.63	7	95.07	96.85
6	91.50	94.36	6	93-33	95.66	6	95.11	96.88
5	91.54	94.38	5	93.37	95.69	5	95.14	96.90
4	91.57	94.41	4	93.41	95.71	4	95.18	96.93
3	91.61	94-43	3	93-44	95-74	3	95.21	96.95
2 1	91.64	94.46	2	93.48	95.76	2	95.25	96.98
6	91.68	94.48	I	93-52	95-79 95-82	I	95.29	97.00
	· · · · / L	74.71	, ,	93-55	1 41.02	. 0	95-32	97.02

SPECIFIC GRAVITY AND PERCENTAGE OF ALCOHOL—(Continued).

	Absolute	Alcohol.	!	Absolute	Alcohol.	1	Absolute	Alcohol.
Spec. Grav at 15.6°C.	Per Cent	Per Cent	Spec. Grav. at 15.6° C.	Per Cent	Per Cent	Spec. Grav. at 15.6° C.	Per Cent	Per Cent
15.0°C.	by Weight.	by Vol- ume.	13.0 C. ¹	by Weight.	by Vol- ume.	13.0 C.	by Weight.	by Vol- ume.
0.8079	95.36	97.05	0.8029	97-07	98.18	0-7979	98.69	99.18
4	95-39	97-07	8	97.10	98.20	8	98.72	99.20
7	95.43	97.10	7 6	97.13	98.22 98.24	7 6	98.75	99.22
6	95.46	97.12 97.15	5	97.16 97.20	98.24 98.27	5	98.78 98.81	99.24
5 4	95.50 95.54	97.17	3 4	97.23	98.29	4	98.84	99.27
3	95-57	97.20	3	97.26	98.31	3	98.87	99.29
2	95.61	97.22	2	97.30	98.33	' ž	98.91	99.31
1	95.64	97.24	il r	97-33	98.35	1	98.94	99.33
0	95.68	97-27	•	97-37	98.37		98.97	99-35
0.8069	95.71	97.29	0.8019	97.40	98.39	0.7969	99.00	99-37
8	95-75	97-32	8	97-43	98.42	8	99.03	99 39
7 6	95.79	97 - 34	7	97.46	98.44	i; 7 6	99.06	99.41
	95.82 95.86	97 - 37	6	97.50	98.46 98.48		90.10	99.43
5 4	95.89	97-39 97-41	5 4	97-53	98.50	5 4	99.13	99-45
3	95.93	97-41	3	97.57	98.52	3	99.10	99.47
2	95.96	97.46	3	97.63	98.54	2	99.23	99.49
I	96.00	97-49	1	97.66	98.56	1	99.26	99.53
•	96.03	97.51	•	97.70	98.59	0	99.29	99.55
0.80 59	96.07	97-53	0.8009	97.73	98.61	0.7959	99.32	99-57
8	96.10	97-55	8	97.76	98.63	8	99.36	99-59
7 6	96.13	97-57	7	97.80	98.65	7	99-39	99.61
	96.16	97.60	6	97.83	98.67	6	99.42	99.63
5	96.20 96.23	97.62 97.64	5	97.87	98.69 98.71	5	99.45	99.65
4 3	96.26	97.66	4 3	97-90 97-93	98.74	4 3	99.48	99.67
2	96.30	97.68	2		98.76	2	99.55	99.09
I	96.33	97.70	1	98.00	98.78	1	99.58	99-73
0	96.37	97-73	0	98.og	98.80	0	99.61	99-75
0.8049	96.40	97-75	0.7999	98.06	98.82	0.7949	99.65	99-77
8	96.43	97-77	8	98.09	98.83	8	99.68	99.80
7	96.46	97-79	7	98.12	98.85	7	99-71	99.82
6	96.50	97.81	6	98.16	98.87	6	99-74	99.84
5	96.53	97.83	5	98.19 98.22	98.89 98.91	5	99.78	99.86
4	96.57 96.60	97.86 97.88	4	98.22	08.93	4	99.81	99.88
3 2	96.63	97.90	3 2	98.28	98.94	3 2	99.87	99.90
1	96.66	97.92	l î	98.31	98.96	ı	99.90	99.92
ō	96.70	97-94	0	98.34	98.98	0	99.94	99.96
0.8039	96.73	97.96	0.7989	98.37	99.00	0.7939	99.97	i . 99.98
⁸	96.76	97.98	8	98.41	99.02		1	1
7	96.80	98.01	7	98.44	99.04		Abs.	Alc.
6	96.83	98.03	6	98.47	99.05	0.7938	100.00	100.00
5	96.87	98.05	5	98.50	99.07			i
4	96.90	98.07	4	98.53	99.09			
3	96.93	98.09	3	98.56	99.11			1
2	96.96	98.11	2	98.59 98.62	99.13]		l
1	97.00	98.16	1 0	98.66	99.15			1
3	97.03	90.10	: 1	90.00	79.10	.1		1

(4) Determination of Alcohol by the Ebullioscope or Vaporimeter is based on the variation in boiling-point of mixtures of alcohol and water, in accordance with the amount of alcohol present. There are various forms of this instrument, one of the simplest and most convenient being that of Salleron, Fig. 113, the apparatus being known in France as an

Fig. 113.—Salleron's Ebullioscope and Scale for Calculation of Results,

ebulliometer. This consists of a jacketed metallic reservoir, heated by a lamp placed beneath, and fitted with a return-flow condenser at the top and with a delicate thermometer graduated in tenths of a degree.

As the boiling-point of water varies with the atmospheric pressure, it is necessary to determine the actual boiling-point corresponding with the barometric conditions each time a series of determinations are made.

This is done by boiling a measured portion of distilled water in the reservoir, and carefully noting the temperature when it becomes constant.

The reservoir is then rinsed out with a little of the liquor to be tested. after which a measured amount of this liquor is boiled in the reservoir and the temperature again noted. A sliding scale (Fig. 113) accompanies the instrument, having three graduated parts as shown. The central movable portion is graduated in degrees and tenths of a degree centigrade, the part at the left has the per cent of alcohol corresponding to the temperature in the case of simple mixtures of alcohol and water. while the part at the right is used for reading the per cent in the case of wine, cider, beer, etc., which have a considerable residue. The movable scale bearing the degrees of temperature is first set with the actual temperature of boiling water (as ascertained) opposite the o mark on the stationary scale. Suppose the temperature of boiling water has been found to be 100.1°. The scale is in this case set as shown in Fig. 113. Suppose also the temperature of boiling of the wine to be tested is found to be 80.3°. From the right-hand scale the corresponding per cent of alcohol is found to be 17.2.

When the liquor to be tested contains more than 25% of alcohol, it is necessary to dilute with a measured amount of distilled water and calculate the per cent from the dilution.

When once the boiling-point of water has been determined for a given barometric pressure, it is unnecessary to change the position of the sliding scale during a series of alcohol determinations unless that pressure changes.

Expression of Results.—Some confusion is caused by the three ways of expressing results of the alcohol determination, whether as per cent by weight, per cent by volume, or grams per 100 cc. The particular mode adopted should depend upon the nature of the case and upon the prevailing custom. In laboratory analyses, unless otherwise qualified, the simple expression of "per cent" usually implies per cent by weight, and for the reason that this conforms with other determinations, the adoption of the weight-percentage plan is perhaps most natural to the chemist on the grounds of uniformity.

In enforcing the laws regulating the liquor traffic, the custom leans to volume percentage, and many of the laws are based on the "volume of alcohol at 60° F." (see p. 656).

In recent years many European analysts have adopted the custom of expressing results of analyses of wines and other liquors in grams per

100 cc. and, in order to have a common basis of comparison between the composition of American and of European wines, this manner of expression has to some extent been adopted in the United States.

Proof-spirit in the United States is an alcoholic liquor containing 50% of absolute alcohol by volume at 15.6° C. A common method of expressing alcohol is in "degree proof" or simply "proof," which in the United States is twice the per cent of alcohol by volume. Thus, 91.3 proof or degree proof is the same as 45.65% alcohol by volume.

English Proof-spirit differs from that in the United States in that it contains 49.24% by weight, or 57.06% by volume of absolute alcohol at 15.6° C. Strength is expressed in degrees over or under proof. Thus liquor 20° under proof has 80 parts by volume of proof-spirit and 20 parts of water at 15.6° C., while 20° under-proof means that 100 volumes of the liquor have to be diluted to 120 volumes with water to yield proof-spirit. To calculate the per cent by volume of English proof-spirit from the per cent of alcohol by volume, divide the latter by 0.5706, or multiply it by 1.7525.

Direct Determination of Extract.—In liquors having a high sugar content, the extract or total solids cannot be determined accurately by evaporation at the temperature of boiling water, owing to the dehydration of the reducing sugars at temperatures exceeding 75°. When extreme accuracy is required, such liquors should be dried *in vacuo* at 75°, or in a McGill oven (p. 586).

Approximate results satisfactory in most cases are obtained by heating for two and one-half hours 10 grams of the liquor in a tared platinum dish at the temperature of boiling water. If the results are to be expressed in grams per 100 cc., instead of weighing out 10 grams, 10 cc. of the liquor are measured by a pipette into a tared dish. With distilled liquors having low residues, accurate results are obtainable by direct evaporation at 100°, using preferably 25 grams or 25 cc. according as the result is to be expressed in per cent by weight or grams per 100 cc.

Extract in wine and beer is more readily calculated indirectly from their specific gravity as noted elsewhere.

Determination of Ash.—The residue from the determination of the extract is incinerated to a white ash in the original dish at a low red heat, either over a Bunsen flame or in a muffle. The dish is finally cooled in a desiccator and weighed.

Preservatives and Artificial Sweeteners in liquors are identified as described in Chapters XVIII and XIX.

FERMENTED LIQUORS.

The fermented juices of many varieties of fruits and berries furnish beverages more or less popular in various localities, especially for home consumption, though, with the exception of the products of the apple and the grape, few of them are found on the market. The following table shows the average percentage of sugar and free acid in the expressed juice or must of fruits, according to Fresenius, arranged in the order of their sugar content:

	Per Cent Sugar.	Per Cent Free Acid as Malic.
Peaches	1.99	0.85
Apricots	2.13	I.20
Plums	2.80	I.72
Green gages	4.18	0.67
Raspberries	4.84	1.80
Blackberries	5-32	1.42
Strawberries	6.89	1.57
Currants.	7.30	2.43
German prunes		r.08
Gooseberries	8.00	1.63
Pears	8.43	0.00
Apples	9.14	0.82
Mulberries	10.00	2.02
Sour cherries	10.44	1.52
Sweet cherries	15.30	o. <u>8</u> 8
Grapes	16.15	0.80

CIDER.

Cider is the expressed juice of the apple. When fresh and before fermentation has set in, it is known as sweet cider, but it does not long remain in this condition, developing after a good fermentation from 3 to 6 per cent of alcohol by volume.

The predominating yeast under the influence of which the fermentation of cider takes place is *Saccharomyces apiculatus*, found in considerable quantity on the outside of the apples as well as in the soil in which the trees grow.

Process of Manufacture.—The best cider is made from ripe fruit, taking care to avoid the green and the rotten apples, both of which impair the quality of the product. After gathering, the apples are best allowed to stand in piles until perfectly ripe, being kept under cover. If exposed to the weather, certain of the yeast organisms found on the skins of the apples that are useful in promoting subsequent fermentation would be

washed off. As a rule the apples commonly used by farmers for cidermaking are those that are unsalable or unfit for other purposes, being chiefly windfalls or bruised and imperfect fruit. The apples are usually first crushed in a mill to a coarse pulp, which is afterward subjected to pressure in a suitable press and the juice thus extracted.

In this country but little attention is paid to the after processes, the juice being usually transferred directly to barrels, which are not always particularly clean, and allowed to ferment spontaneously in a convenient place, subject to changes in temperature. There is little wonder that cider so made will keep but a short time and quickly goes over into vinegar, unless salicylic acid or other antiseptic is added.

In France more care is taken to regulate the temperature of fermentation, to insure absolute cleanness of all receptacles, and to separate out contaminating impurities. A preliminary fermentation is usually given to the juice in open vats, during which the yeast spores are developed, while impurities separate out both by rising to the surface and by settling to the bottom, care being taken to avoid the development of acetic fermentation. At the proper time the juice is "racked off" or drawn from the clear portion between the top and bottom, transferred to scrupulously clean barrels, and allowed to undergo a second fermentation at a lower temperature than before.

Sometimes the "racking off" is repeated, and the juice is further clarified by "fining" or treating with isinglass, which carries down certain albuminous substances.

Cider thus made is capable of keeping a very long time.

In England cider is sometimes "fined" by treatment with milk, one quart of the latter being added to eighteen gallons of cider.

The apple pomace, left as a residue, is generally steeped in water and repressed. The juice from the second pressing is occasionally added to the first for cider manufacture, but more often is concentrated and made into apple jelly, or used as a fortifier for vinegar to make up deficiency in solids.

Composition of Cider.—The following tables, due to Browne,* show the chemical composition of the freshly expressed juice of several American varieties of apple, as well as that of a few fermented samples of cider of known purity.

^{*} Penn. Dept. of Agric., Bul. 58.

APPLE JUICES.

	Specific Gravity.	Solids.	Invert Sugar.	Sucrose.	Total Sugar.	Total Sugar after Inversion.	Free Malic Acid.	Ash.	Undetermined, Pectin, etc.	Rotation, 400-mm. Tube, Ventzke Scale. Degrees to the Left.
Red astrachan Early harvest. Yellow transparent Early strawberry. Sweet bough Baldwin, green. "ripe. Ben Davis. Bellflower. Tulpahocken. Unknown.	1.05317 1.05522 1.05020 1.04049 1.04079 1.04882 1.07362 1.05389 1.06270 1.05727	11.71 11.81 11.87 11.36 16.82 12.77 14.90	7-49 8-03 5-47 7-61 6-96 7-97 7-11 9-06 9-68	3-97 2.10 4.21 3.08 1.63 7.05 3.85 4-32 3.11	11.46 10.14 9.68 10.69 8.59 15.02 10.96 13.38	10.60 11.67 10.24 9.00 10.85 8.68 15.39 11.16 13.61	0.90 0.86 0.78 0.10 1.24 0.67 0.46 0.58 0.26	0.28 0.27 0.24 0.31 0.26 0.28 0.28	0.65 0.44 1.11 1.22 0.87 1.07 0.66 0.49	23.72 24.32 19.24 39.40 36.16 49.00 39.20 48.20 44.18

FERMENTED CIDER (MIXED APPLES).

Spec Grav		Solids.	Invert Sugar.	Malic Acid.	Acetic Acid.	Alcohol.	Pectin.	Ash.	Rotation, 400-mm. Tube, Ventzke Scale. Degrees to the Left.
A 1.99	805	1.94	0.19	0.21	0.24	6.85	0.03	0.25	2.30
B 1.00	122	2.71	0.19	0.24	0.42	5.13	0.03	0.32	2.49
C 1.00	525	3.26	0.89	0.30	0.48	4.67	0.05	0.29	5.28
D 1.00	071	1.93	0.34	0.27	0.21	4-95	0.05	0.23	2.00
E 1.00	512	2.71	0.24	0.29	1.96	4.26	0.06	0.36	1.76

The following are summaries of the results of a large number of analyses of European apple juices made by Truelle, the quantities being expressed in grams per liter:

	Mean.	Minimum.	Maximum.
Specific gravity	1.0760	1.0573	1.1100
Inverguagar	135.85	108.38	181.81
Sucrose	25.01	5.58	71.7
Total fermentable sugars (as dextrose)	162.18	119.22	231-57 8.07
Tannin	2.90	0.26	8.07
Pectin and albuminous substances	12	0	23
Acidity (sulphuric acid)	2.14	0.69	7-41

In the municipal laboratory of Paris, Sanglé Ferrière has analyzed eleven samples of known-purity cider with the following results:

				Suge Li	r per ter.			Ļ	Acidi H ₂	ity as
	Density.	Per Cent Alcohol by Volume.	Extract per	Before Inver- ston.	After Inver-	Polarization Laurent,	Ash per Liter.	Ash, as XxCO, per Later.	Total.	Fixed
Mean Maximum Minimum	1.0159 1.0410 1.0012	3.9 6.2 1.1	52.67 114.00 22.62	21.31 59.40 Trace	21.62 60.80 Trace	-4°.26 -11°.20 0	3.26 4.32 2.48	2.56 3.68 2.04	5.27 6.59 4.20	2.55 2.94 1.47

Six samples of bottled "sweet" cider purchased in Massachusetts were analyzed in the Food and Drug Laboratory of the Board of Health with the following results:

	Per Cent Alcohol by Weight.	Per Cent Acid as Malic.	Per Cent Extract.
Maximum Minimum Average	3-55	0-72 0-48 0-58	7.82 2-42 4-19

Browne gives the following as the composition of the mixed ash of several varieties of apple:

Potash (K ₂ O) Soda (Na ₂ O) Lime (CaO). Magnesia (MgO). Oxide of iron (Fe ₂ O ₂). Oxide of aluminum (Al ₂ O ₃). Chlorine (Cl). Silica (SiO ₂). Sulphuric acid (SO ₁). Phosphoric acid (P ₂ O ₃). Carbonic acid (CO ₂).	4-43 3-78 0-95 0-80 0-39 0-40 2-66 8-64	Equivalent to	Potassium carbonate (K ₂ CO ₃) Potassium phosphate (K ₂ PO ₄) Sodium chloride (NaCl) Calcium sulphate (CaSO ₄) Calcium oxide (CaO). Magnesium phosphate (Mg ₂ P ₂ O ₄) Magnesium oxide (MgO) Ferric oxide (Fe ₂ O ₃) Aluminum oxide (Al ₂ O ₃) Silica (SiO ₂)	6.85 14-55 0.60 4-5 ² 2-57 6.97 0.59 0.95 0.40
Deduct oxygen equivalent to Cl Total	99.90 .09 99.81		Total	99.80

Burcker * gives the following composition of the ash of cider:

	Per Cent.
Silica	0.94
Phosphoric acid	12.68
Lime	2.77
Magnesia	2.05
Oxides of iron and manganese	0.94
Potash	53-74
Soda	1.10
Carbonic acid	25.78
•	
	00.00

Adulteration of Cider.—The Committee on Standards of the A. O. A. C. have submitted for adoption the following standards for cider: Alcohol not more than 8%, extract not less than 1.8% determined by evaporation in an open vessel at ordinary atmospheric pressure and at the temperature of boiling water; ash not less than 0.2%.

Entirely factitious cider made from other than apple stock is rarely found, though the product as sold is frequently of inferior quality and adulterated. The chief adulterants are water and sugar, and the use of antiseptics is common, especially of salicylic and sulphurous acids, sodium benzoate, and occasionally beta-naphthol.

Sodium carbonate is sometimes added to cider to neutralize the acid and thus prevent acetic fermentation. An abnormally high ash (say in excess of 0.35%) would point toward the presence of added alkali.

Watering is apparent when the content of alcohol, solids, and ash of the suspected sample are found to be considerably below the corresponding constants of pure cider. According to Sanglé Ferrière, the following are the minimum figures for these constants in a pure cider, so that a sample may safely be pronounced as watered if they all run distinctly below:

Alcohol	3% by volume
Extract	1.8%
Ash	0.17%

Besides these determinations, it is useful also to determine the fixed and volatile acids.

Caramel is to be looked for, especially in watered samples. Other

^{*} Les Falsifications des Substances Alimentaires, p. 176.

adulterants alleged to be of frequent occurrence in French cider, but not commonly found in this country are commercial glucose, tartaric acid (to increase the acidity of a watered product), and coal-tar colors.

Absence or deficiency of malates is conclusive evidence of fraud, indicating the admixture of notable quantities of the juice of the second pressing of pomace.

Sugar is rendered apparent by the right-handed polarization of the sample, pure cider always polarizing well to the left. If after inversion of a dextro-rotary cider the polarization is still to the right, commercial glucose is indicated; if the reading after inversion is to the left, cane sugar has undoubtedly been added.

Frequently the analyst has only to determine the alcohol, especially in cases of seizure, to ascertain whether or not there has been violation of the liquor laws.

PERRY OR PEAR CIDER.

This is a common French product, but is rarely if ever found on sale in this country, though sometimes made for home consumption. In composition and in method of manufacture it much resembles apple cider. It is also subject to the same forms of adulteration.

The following table summarizes a number of analyses made by Truelle on pear juice, or must, amounts being expressed in parts per thousand:

	Mean.	Maximum.	Minimum.
Specific gravity.	1.0845	1.0675	1.0980
Invert sugar. Sucrose	145.64 36.74	108.10	200 61.41
Total fermentable sugars (as dextrose)	184.14	143.78	220
Tannin	1.78	1.01	3.20 18
Pectin and albuminous substances Acidity (as sulphuric acid)	13.08 1.47	3 0.76	18 2.40

The following analysis of champagne perry is taken from the *Lancet* of October 1, 1892:

Alcohol by weight	1.45
Alcohol by volume	1.80
Solids	11.00
Ash	0.25

WINE.

Wine in its broadest sense is the fermented expressed juice of any fruit, though the term, unless otherwise restricted, is generally understood to apply to the juice of the grape.

The organism present in grape juice that plays the chief part in its alcoholic fermentation is the *Saccharomyces ellipsoideus*, a yeast which exists on the skins of the grape.

Process of Manufacture.—The grapes, which should be fully ripe, are picked and sometimes sorted, according to the care that is taken in grading the product. They are also sometimes freed from the stems, which contain considerable tannic acid, and which when crushed with the grapes impart a certain astringency to the final product. The grapes are crushed either by machinery or by the bare feet, and the juice is pressed out from the pulp in various ways, by screw or hydraulic press, or by the centrifugal process.

A certain amount of juice runs off from the preliminary crushing known as the first run, and makes the choicest wine. The product from the pressure constitutes the second run, after which the pomace, by steeping in water and repressing, is made to yield an inferior juice used in vinegar-making.

Red wines are made from dark grapes by fermenting the pulp, beforepressing, with the skins, which by this treatment yield up their rich color (*anocyanin*) to the juice. Besides the color, the skins contain also tannin. White wine is made from the pressed pulp, freed from the skins at once, or from the pulp of white grapes. The unfermented must constitutes from 60 to 80 per cent of the weight of the grape.

Fermentation progresses most rapidly at a temperature between 25° and 30° C., but wine having a much finer bouquet is produced by slower fermentation, hence the must is allowed to ferment in open vats or tubs in cool cellars, at a temperature of from 5° to 15° till it settles out comparatively clear, special care being taken to avoid development of acetic fermentation. At the end of the first or active fermentation, the wine is drawn off and allowed to undergo a second or slow fermentation in casks, during which most of the lees or crude argols, composed of potassium bitartrate, settle out, being insoluble in alcohol, and the characteristic bouquet or flavor of the wine is developed. Occasionally during this process the wine is racked or drawn off.

Undesirable fermentations and vegetable fungus growth, which are

liable to occur at this time, are avoided as much as possible by using especially clean casks, which are frequently "sulphured" (or burnt out with sulphur) before being used. The wine is also sometimes clarified, or "fined," by treatment with gelatin, which mechanically removes many impurities by precipitation, or is subjected to pasteurization before finally being bottled or stored in casks.

Classification of Wines.—Wines are either natural or jortified. Natural wines are those which contain no added sugar or alcohol, but which are exclusively the product of the simple juice, fermented under the best conditions, either till the sugar has been used up, or till the yeast food is exhausted, or until the yeast growth has been checked by the strength of the alcohol developed. When the alcohol content amounts to 14% by weight there can be no further fermentation due to yeast, so that this is the highest limit for natural wine. Examples of natural wines are hock and claret and many California wines.

Fortified wines are those to which alcohol has been added, usually before the natural fermentation has been allowed to proceed to a finish. For this reason considerable sugar is usually left, and such wines are more often sweet. Examples of fortified wines are Madeira, sherry, and port.

Volatile ethers (products of volatile acids) predominate as a rule in natural wines, while fixed ethers (from the fixed acids as tartaric) are most characteristic in fortified wines.

Wines are also variously classified according to characteristic properties possessed by them, as still or sparkling, red or white, "dry" or sweet, etc.

Still wines are those in which there is but little carbon dioxide remaining, so that they do not effervesce. Sparkling wines are more or less heavily charged with carbon dioxide, either naturally, as in the case of champagne, wherein the gas is formed by after-fermentation of added sugar in the corked bottle, or artificially, by carbonating them in a similar manner to "soda-water."

Among the best-known red wines are those of Burgundy and the Bordeaux wines or clarets, while the Rhenish and Moselle wines and the Sauternes are examples of white wine.

"Dry" wines are those in which the sugar has been exhausted by fermentation, while sweet wines possess a considerable amount of unfermented sugar. Whether or not an excess of sugar is left after fermentation has stopped depends upon the amount of yeast food or nitrogenous substance present in the wine. When the proteins are exhausted by the yeast, fermentation ceases, and for this reason gelatin and other nitrogenous bodies are sometimes added to extend the period of fermentation. Sweet wines are often reinforced by the addition of sugar. Madeira, both red and white, are samples of dry wine, while port wine is one of the sweet variety.

While most of our finer wines still come from France and Germany, large quantities of California wines are now being produced of an extremely high grade and of many varieties.

Composition of Grape Must and of Wine.—König's analyses of a large number of grape musts from different sources are thus summarized:

	Specific Gravity.	Water, Per Cent.	Nitroge- nous Ma- terial.	Sugar.	Acid.	Other Non-ni- trogenous Material.	Ash.
Minimum	1.0690	51.53	0.11	12.89	0.20	1.68	0.20
	1.2075	82.10	0.57	35-45	1.18	11.62	0.63
	1.1024	74.49	0.28	19.71	0.64	4.48	0.40

Typical analyses of German, French, Austrian, Russian, Italian, and Spanish wines are shown in the following table, also due to König:

	Number of Analyses.	Specific Gravity.	Alcohol by Weight.	Extract.	Total Acid as Tartaric.	Free Tar- taric Acid.	Cream of Tartar.	Volatile Acid as Acetic.	Sugar.
Germany:					-				
Moselle	14	0.9964	7.99	2.24	0.79				0.031
Rhine	23	1.0005	8.00	2.60	0.81		0.20		
Baden	46		6.65	2.16	0.91	0.018	0.358		0.005
Wurtemburg, white wine.	15	0.9995	6.10	2.27	0.95	0.095			
" red wine	6		4-73	2.64	1.14	0.091			
Alsace	15		6.59	2.07	0.696	0.018	0.168	0.052	
Lorraine, red wine	10	0.9967	8.08	2.27	0.56	0.032		0.155	0.088
France:									
Red wine	29	0.9982	7.80	2.56	0.57				0.30
White wine	5	0.9963	8.30	3.03	0.66				_
Austria:		İ							
Tyrol, red wine		0.9940		2.34	0.62		- 1		
" white wine	17	0.9927	8.84	1.87	0.59				
Russia:							1	-	
Red wine		0.9939		2.76	0.56			0.142	
White wine		0.9931		2.568	0.49			0.100	0.458
Italy	20		10.61	3-44	0.52				1.44
Spain:							i		
Ordinary red wine	7		I	3-53	0.49				0.38
Sweer wine	4	1.0233	12.78	9.69	0.59				6.55

	Glycerin.	Nitrogen.	Ash.	P ₈ O ₆ .	Potash.	Lime.	Magnesia.	Sulphuric Acid.	Chlorine.
Germany: Moselle Rhine. Baden	0.85		0.23	0.046	0.085	0.011		0.012	
Wurtemburg, white wine. " red wine. Alsace	0.57 0.40 0.55	0.028	0.25 0.25 0.229	0.043 0.040 0.038	0.115	· ·	0.010	0.08	0.023
France: Red wine White wine	0.73	0.043	0.248	0.030	0.106	0.101		0.033	
Austria: Tyrol, red wine '' white wine Russia:									
Red wine	0.59	0.026	0.204		0.086		0.017	0.019	
Ordinary red wine Sweet wine				0.027 0.039			•••••		

On page 688 are given summaries of analyses of American wines compiled from tables of analyses made by Bigelow.*

Varieties of Wine.—Champagne is a selected, sweet, white wine, clarified with gelatin, bottled with the addition of cane sugar, mixed with a little brandy, and tightly corked. Sometimes a small amount of yeast is also introduced. Fermentation is allowed to go on at a temperature of about 24° C., during which the wine is highly charged with carbon dioxide. The bottles are set on their side for some months, after which they are inverted till the sediment gathers above the cork, which by careful manipulation is quickly removed so as to throw out the sediment, and is afterward replaced and secured. Champagne contains from 8 to 10 per cent of alcohol and is high in sugar.

Claret is a light, red wine of a deep color, and is somewhat acid and astringent. In alcohol it varies from 8 to 13 per cent by volume. It has very little sugar and is high in volatile ethers.

Madeira is a strong, white wine, possessing a refined, nutty, aromatic flavor when fully aged. It is generally fortified, containing from 17 to 20 per cent of alcohol. It is named from the island which produces it. Sherry is a deep, amber-colored, sweet, Spanish wine, high in alcohol.

^{*} U. S. Dept. of Agric., Bur. of Chem., Bul. 59.

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	Ash.	200.		.416	. 202	.140	.368	. 290 . 148	.394	.436
g	Tennin and Coloring Matter.	.558	.349	.328	.050		.015	.045	1.066	.350
Grams per 100	Proteids.	.5544 .1865		.1864		.5164	.38%	.3162	.9379 .1893	.0859
Gram	muissatoq Sulphate.	.1570		.2515		.0633	.1648		.0594	.0504
	Reducing Sugar.	.040		.030		.626	3.569	.936	13.559	23.117.210
	noitazitalo¶	-2.1	: :	-1.7 -0.5		13.5	+0.6 -18.6	-3.4 -0.1	-27.1 -14.4	-23.1 -0.5
Grams per	Extract.	3.81	3.34	3.46	6.88*	4.38	6.78	4.56	17.22	19.66
Gram 100	Total Acids.	.368	.358	.762 .408	.834	.788	.766	.219	.780 .181	.253
	Glycerin- alcohol Ratio.	8.7:100		7:100		10.2:100	7.7:100	3.4:100	4.5:100	6.8:100
per cc.	Glycerin.	.852	::	.656		.971	.904	.318	.707	.936
Grams per	Alcohol.	7.00	10.96	6.35	15.30	3.98	6.53	17.60	17.61	17.34
	Per Cent Alcohol by Volume,	15.09	13.82	15.48	19.28*	14.57	15.20	8.07	22.19 10.38	8.22
	Specific Gravity.	1.0020	-9970	.9962	1.0050 .9900	1.0020	1.0157	.9988	1.0429	1.0560
	Number of Samples.	8r 6	2	3	\	87	3 : :3	, ; ; 0	, ; ; %	
		RED WINES. Bordeaux, or claret type Maximum. Minimum.	type	Maximum. Minimum. Southern French tyne		WHITE WINES. Rhine-wine type. Maximum. Minimum. Sauteme type.	Maximum. Minimum.		Maximum. Minimum.	

ortified.

(sometimes containing over 20%), being usually fortified. It is slightly acid and possesses much fragrance. Sherry is nearly always "plastered."

Hocks are white German wines, mildly acid, containing 9 to 12 per cent of alcohol by volume. They have very little sugar, and rank among the highest of natural wines. The best-known varieties are Hockheimer and Johanisberger.

Port (Vinum portense of the 1870 Pharmacopæia) is a dark-purple, astringent wine, almost always fortified, and hence high in alcohol (from 15 to 18 per cent by volume). It is much improved by aging, during which it looses considerable of its astringency. It contains a large amount of extract, from 2 to 6 per cent of the wine being sugar. The fixed ethers predominate over the volatile.

Standards of Purity for Wine.—The ratio of volatile to fixed acids in pure wine should not exceed 1:3. A higher proportion of volatile acid shows the fact that acetic fermentation has set in.

The presence of any considerable free tartaric acid would indicate the addition of this substance to the wine.

The United States Pharmacopæia has prescribed the following requirements in the case of wines: For white wine (Vinum album) the specific gravity at 15.6° should not be less than 0.990 nor more than 1.010; the extract or residue at 100° should not be less than 1.5 nor more than 3%; as indicating the amount of free acid, not less than 3 nor more than 5.2 cc. normal potassium hydroxide should be required to neutralize 50 cc. of the wine, using phenolphthalein as an indicator; it should contain not less than 7 nor more than 12 per cent by weight of absolute alcohol; it should contain only traces of tannin.

For red wine (Vinum rubum) the specific gravity at 15.6° should not be less than 0.989 nor more than 1.010; the extract should not be less than 1.6% nor more than 3.5%; its limits as to acidity are the same as with white wine, eosin or fluorescin, however, being used as an indicator; in alcoholic strength, it should, like white wine, come within the limits of 7 and 12 per cent alcohol by weight. It should not be artificially colored, but should show the presence of tannic acid.

The following are U. S. standards for wines: Wine is the product made by the normal alcoholic fermentation of the juice of sound, ripe grapes, and the usual cellar treatment, and contains not less than 7 nor more than 16 per cent of alcohol, by volume, and, in 100 cc. (20° C.),

not more than 0.1 gram of sodium chloride nor more than 0.2 gram of potassium sulphate; and for red wine not more than 0.14 gram, and for white wine not more than 0.12 gram of volatile acids produced by fermentation and calculated as acetic acid. Red wine is wine containing the red coloring matter of the skins of grape. White wine is wine made from white grapes or the expressed fresh juice of other grapes.

Dry wine is wine in which the fermentation of the sugars is practically complete, and which contains, in 100 cc. (20° C.), less than 1 gram of sugars, and for dry red wine not less than 0.16 gram of grape ash and not less than 1.6 grams of sugar-free grape solids, and for dry white wine not less than 0.13 gram of grape ash and not less than 1.4 grams of sugar-free grape solids.

Fortified dry wine is dry wine to which brandy has been added, but which conforms in all other particulars to the standard of dry wine.

Sweet wine is wine in which the alcoholic fermentation has been arrested, and which contains, in 100 cc. (20° C.), not less than 1 gram of sugars, and for sweet red wine not less than 0.16 gram of grape ash, and for sweet white wine not less than 0.13 gram of grape ash.

Fortified sweet wine is sweet wine to which wine spirits have been added. By act of Congress, "sweet wine" used for making fortified sweet wine and "wine spirits" used for such fortification are defined as follows (sec. 43, Act. of October 1, 1800, 26 Stat. 567, as amended by section 68, Act of August 27, 1894, 28 Stat. 509, and further amended by Act of Congress, approved June 7, 1906): "That the wine spirits mentioned in section 42 of this act is the product resulting from the distillation of fermented grape juice to which water may have been added, prior to, during, or after fermentation, for the sole purpose of facilitating the fermentation, and economical distillation thereof, and shall be held to include the products from grapes or their residues, commonly known as grape brandy; and the pure sweet wine, which may be fortified free of tax, as provided in said section, is fermented grape juice only, and shall contain no other substance whatever introduced before, at the time of, or after fermentation, except as herein expressly provided; and such sweet wine shall contain not less than 4 per cent of saccharine matter, which saccharine strength may be determined

by testing with Balling's saccharometer or must scale, such sweet wine, after the evaporation of the spirits contained therein, and restoring the sample tested to original volume by addition of water: Provided, That the addition of pure boiled or condensed grape must, or pure crystallized cane or beet sugar or pure anhydrous sugar to the pure grape juice aforesaid, or the fermented product of such grape juice prior to the fortification provided by this act, for the sole purpose of perfecting sweet wine according to commercial standard, or the addition of water in such quantities only as may be necessary in the mechanical operation of grape conveyors, crushers, and pipes leading to fermenting tanks, shall not be excluded by the definition of pure sweet wine aforesaid: Provided, however. That the cane or beet sugar, or pure anhydrous sugar. or water, so used shall not in either case be in excess of 10% of the weight of the wine to be fortified under this act: And provided further, That the addition of water herein authorized shall be under such regulations and limitations as the Commissioner of Internal Revenue, with the approval of the Secretary of the Treasury, may from time to time prescribe; but in no case shall such wines to which water has been added be eligible for fortification under the provisions of this act where the same, after fermentation and before fortification, have an alcoholic strength of less than 5% of their volume."

Sparkling wine is wine in which the after part of the fermentation is completed in the bottle, the sediment being disgorged and its place supplied by wine or sugar liquor, and which contains in 100 cc. (20° C.), not less than 0.12 gram of grape ash.

Modified wine, ameliorated wine, corrected wine, is the product made by the alcoholic fermentation, with the usual cellar treatment, of a mixture of the juice of sound, ripe grapes with sugar (sucrose), or a syrup containing not less than 65% of sugar (sucrose), and in quantity not more than enough to raise the alcoholic strength after fermentation, to 11% by volume.

Raisin wine is the product made by the alcoholic fermentation of an infusion of dried or evaporated grapes, or of a mixture of such infusion, or of raisins with grape juice.

Adulteration of Wine.—Beverages purporting to be wine are sometimes found on sale that are entirely spurious, in that they contain little if any fermented grape juice. Apple cider is not infrequently the basis of such artificial products, and the following recipes given

by Brannt may be taken as typical of the composition of these wine substitutes:

Burgundy.—Bring into a barrel 40 quarts of apple juice, 5 pounds of bruised raisins, \(\frac{1}{4} \) pound of tartar, I quart of bilberry juice, and 3 pounds sugar. Allow the whole to ferment, filling constantly up with cider. Then clarify with isinglass, add about I ounce of essence of bitter almonds, and after a few weeks draw off into bottles.

Malaga Wine.—Apple juice, 40 quarts; crushed raisins, 10 pounds; rectified alcohol, 2 quarts; sugar solution, 2 quarts; elderberry flowers, 1 quart; acetic ether, 1 ounce and 2 drachms. The desired coloration is effected by the addition of bilberry or elderberry juice; otherwise the process is the same as given for Burgundy.

Sherry Wine.—Apple juice, 50 quarts; orange-flower water, about 2 drachms; tartar, 2 ounces and 4 drachms; rectified alcohol, 3 quarts; crushed raisins, 10 pounds; acetic ether, 1 ounce and 2 drachms. The process is the same as for Burgundy.

Claret Wine.—Apple juice, 50 quarts; rectified alcohol, 4 quarts; black currant juice, 2 quarts; tartar, 2 ounces and 4 drachms. Color with bilberry juice. The further process is the same as for Burgundy.

Artificial products similar in nature to the above are also mixed in varying proportions with pure wine.

Presence of malates, as well as absence or diminution of total tartaric acid. is also indicative of cider.

If the ash of the wine be submitted to the flame test, the sodium light will predominate in the case of pure wine, while if the basis of the sample be largely or wholly apple stock, the potash flame will be readily apparent.

Wines are most frequently adulterated by "plastering," by watering, by the addition of excessive amounts of sugar or glucose, by various flavoring principles, by coal-tar and vegetable colors, by antiseptics, and by added alcohol.

Plastering.—By this term is understood the addition of gypsum or plaster of Paris to the must before fermentation, a practice in vogue in parts of France, Italy, and Spain. The reaction which takes place with the potassium bitartrate present in the wine is, according to Chancel, as follows:

Various advantages are said to result from this practice. The wine is clarified by the precipitation of the calcium tartrate, which mechanically carries down with it many impurities, the color of the wine is improved, since the solubility of the coloring principle present in the skins is increased, the fermentation is rendered more rapid and complete, and the keeping qualities of the wine are enhanced. The practice is, however, considered objectionable on account of the potassium sulphate which is left in solution in the wine, and in some countries plastering is forbidden, or the amount of potassium sulphate limited by statute.

The following are analyses of two Spanish wines made from the same grape juice, one of which was plastered. The results are expressed in grams per liter.

Not Plastered.	Plastered.
Yellow	Bright red
	27·3 0.61
2.06	5.38
1.29	0.17
0.41	5
	Yellow 23.3 0.66 2.06

The effect of plastering is thus seen to distinctly increase the extract and the soluble ash. Any considerable amount of potassium sulphate is an indication of plastering.

Addition of Cane Sugar.—The term "chaptalizing" is applied in France to the addition to the must of cane sugar for the purpose of increasing the yield in alcohol. The addition of 1,700 grams of sugar to 1,000 liters of must is said to increase the alcoholic strength by 1%. It was formerly customary to add with the sugar calcium carbonate, to partially neutralize the acidity, but this is rarely practiced at present.

The European wine-raising countries are not disposed to regard the reinforcement of wine by added cane sugar in the must as in itself a fraud, unless water is also added. or unless some other form of adulteration is practiced at the same time. In France, however, the addition of cane

sugar is permitted only in wine for local consumption, and is restricted in amount.

The use of commercial glucose in wine instead of cane sugar is not regarded with as much favor, in view of the fact that glucose contains more or less unfermentable matter, and introduces dextrin and various mineral salts into the wine.

To ascertain the nature and extent of the sugars present in wine is frequently of great importance. Much information may be gained from the direct and invert polarization of the sample, as well as from the determination of reducing sugars.

Invert sugar is the only legitimate sugar that should be present in genuine wine. In normal fermentation the dextrose is more quickly destroyed than the levulose, hence the polarization of pure wine is always left-handed, unless all the sugar has been fermented, in which case the reading should be zero.

Seventy-five samples of California red wines, chiefly claret, Burgundy, Rhine, and southern France types, analyzed in the Bureau of Chemistry * of the U. S. Department of Agriculture, polarized from -0.5 to -2.1. Upwards of eighty samples of California white wine (of the types of Burgundy, Sauterne, and southern France) were submitted to polarization and all but four were left-handed. These four (evidently abnormal) polarized from 0.1 to 0.1 to 0.1 Most of them varied from 0.1 to

Thirteen of the port wine types (California) had a left-handed polarization of from -14.7 to -27.1. These apparently contained large quantities of unfermented, inverted cane sugar.

A sharp, right-handed polarization would indicate the presence of either commercial glucose or cane sugar. After inversion, if the reading is still right-handed, glucose is apparent, while if inversion changes the reading from right to left, cane sugar has undoubtedly been added. By application of Clerget's formula the amount of cane sugar can be estimated.

The Watering of Wine, unless excessive in degree, is not always easy to prove, by reason of the varying composition of pure wine, and because the practice is usually accompanied by other forms of sophistication intended to cover up evidences of watering. Considerable quantities of added water alone would usually be rendered apparent by a proportionate and abnormal lowering of the alcohol, extract, ash, acidity, and, indeed, nearly all the constants.

Gautier in his Traité sur la Sophistication et l'Analyse des Vins claims

that the sum of the weight in grams of alcohol in 100 cc. and the total acidity, expressed in grams of sulphuric acid per liter, varies within very narrow limits in pure wines, rarely being below 13 or above 17. A large number of analyses made by Gautier and others would seem to confirm this, so that in the majority of cases, added water would be strongly indicated if the sum of these two constants was materially reduced below 13. It is more conservative to adopt 12.5 as a minimum limit for the sum of the alcohol and total acid expressed as above.

Detection of Added Alcohol.—As a result of the findings of a committee appointed in France to determine the matter of added alcohol, it was submitted that a relation existed between the weight of the extract and that of the alcohol in pure wine. In the case of red wines, if the weight of the alcohol, divided by the weight of the extract (both expressed in grams per 100 cc.) exceeds 4.6, the addition of alcohol is strongly indicated. With white wines, the quotient obtained by dividing the weight of alcohol by weight of extract should not exceed 6.6. If it does, added alcohol is to be suspected.

In the case of plastered wines containing sulphate of potassium, or wines having added sugar, it is necessary to deduct from the total extract the weight of the reducing sugar and of the potassium sulphate as found (less o. I gram for each of these substances), the difference, or reduced extract as it is called, being used in this case in obtaining the ratio.

Fruit Wines other than Grape.—Wines mostly of domestic manufacture are sometimes made from small fruits, such as raspberries, strawberries, blackberries, gooseberries, elderberries, and currants, as well as from cherries, plums, and apricots. Wines made from most of these fruits readily undergo acetic fermentation unless antiseptics are added, or unless extreme care is taken in their manufacture and keeping. Frequently mixtures of various fruit juices are made to yield excellent wine. Most of the sour fruits require a liberal admixture of sugar to produce an acceptable wine.

The alcoholic content of other fruit wines is thus shown by Brannt:

METHODS OF ANALYSIS OF WINE AND CIDER.

For determination of specific gravity, alcohol, extract (by direct method), and ash, see pp. 657-676.

Calculation of the Extract in Wine.—Attention has already been called to the difficulty in accurately determining the extract of sweet wines gravimetrically by evaporation. An approximate determination of the extract may be obtained by calculation from the specific gravity of the dealcoholized liquor, or one may use for this purpose the tables compiled by Windisch, and based on experiments made on drying wine *in vacuo* at 75° C. In wines high in sugar, with more than 6% of extract, this method is far more accurate than drying at 100°, and is to be recommended.

Evaporate a measured portion of the wine on the water-bath to one-fourth its volume, and dilute with water to exactly the volume measured. Determine the specific gravity of this dealcoholized liquid at 15.6°, and from the following table ascertain the extract corresponding.

Determination of Total Acidity.—Carbonated beverages are first freed from carbon dioxide by agitation as described on page 658, after which 25 cc. of the sample are heated just to the boiling-point and titrated with tenth-normal sodium hydroxide, using in the case of white wine or cider phenolphthalein as an indicator. With red wine delicate litmuspaper should be used. Total acidity is usually expressed, in the case of cider as malic, and of wine as tartaric acid. Each cubic centimeter of tenth-normal alkali corresponds to 0.0067 gram malic, or 0.0075 gram tartaric acid. Some chemists express total acidity in terms of sulphuric acid, each cubic centimeter of tenth-normal alkali being equivalent to 0.0049 gram of sulphuric acid.

Volatile Acids in all liquors are usually expressed as acetic, although traces of propionic and other volatile acids may be present. 50 cc. of the cider or wine and a little tannic acid are transferred to a distilling-flask, Fig. 114, the stopper of which is provided with two tubes, one of which connects with the condenser, while the other, arranged to reach nearly to the bottom of the distilling-flask, communicates with a second flask which contains about 300 cc. of water. The contents of both flasks are brought to boiling, after which the flame under the distilling-flask is lowered, and steam from the water-flask is passed through the wine till about 200 cc. of distillate have collected in the receiving-flask. Titrate this with tenth-normal sodium hydroxide, using phenolphthalein

EXTRACT IN WINE. [According to Windisch.]

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Specific Gravity.	Ex- tract.	Specific Gravity.	Ex- tract.	Specific Gravity.	Ex- tract.	Specific Gravity.	Ex- tract.	Specific Gravity.	Ex- tract.	Specific Gravity.	Ex- tract.
1.0000	0.00	1.0065	1.68	1.0130	3.36	1.0195	5.04	1.0260	6.72	1.0325	8.40
1.0001	0.03	1.0066	1.70	1.0131	3.38	1.0196	5.06	1.0261	6.75	1.0326	8.43
1.0002	0.05	1.0067	1.73	1.0132	3.41	1.0197	5.09	1.0262	6.77	1.0327	8.40
1.0003	o.o8	1.0068	1.76	1.0133	3.43	1.0198	5.11	1.0263	6.80	1.0328	8.48
1.0004	0.10	1.0069	1.78	1.0134	3.46	1.0199	5.14	1.0264	6.82	1.0329	8.51
1.0005	0.13	1.0070	1.81	1.0135	3.49	1.0200	5.17	1.0265	6.85	1.0330	8.53
1.0006	0.15	1.0071	1.83	1.0136	3.51	1.0201	5.19	1.0266	6.88	1.0331	8.56
1.0007	0.18	1.0072	T.86	1.0137	3.54	1.0202	5.22	1.0267	6.90	1.0332	8.59
1.0008	0.20	1.0073		1.0138	3.56	1.0203	5.25	1.0268	6.93	1.0333	8.61
1.0009	0.23	1.0074	1.91	1.0139	3.59	1.0204	5.27	1.0269	6.95	1.0334	8.64
		l i		!				1			
1.0010	0.26	1.0075	1.94	1.0140	3.62	1.0205	5.30	1.0270	6.98	1.0335	8.66
1.0011	0.28	1.0076	1.96	1.0141	3.64	1.0206		1.0271	7.01	1.0336	8.69 8.72
1.0013	0.34	1.0078	1.99	1.0143	3.69	1.0208	5.35 5.38	1.0272	7.03	1.0337	8.74
1.0014	0.36	1.0079	2.04	1.0144	3.72	1.0209	5.40	1.0274	7.08	1.0339	8.77
			1		J			1.33,4	,	1557	
1.0015	0.39	1.0080	2.07	1.0145	3.75	1.0210	5.43	1.0275	7.11	1.0340	8.79
1.0016	0.41	1.0081	2.00	1.0146	3.77	1.0211	5.45	1.0276	7.13	1.0341	8.82
1.0017	0.44	1.0082	2.12	1.0147	3.80			1.0277	7.16	1.0342	8.85
1.0018	0.49	1.0084		1.0140	3.85	1.0213	5.51	1.0278	7.19 7.21	1.0343	8.90
2.50.9	J. 49	1.5004	<i>,</i>		3.03		3.33	1.52/9	7.21	1.0344	J. 90
1.0020	0.52	1.0085	2.19	1.0150	3.87	1.0215	5.56	1.0280	7.24	1.0345	8.92
1.0021	0.54	1.0086	2.22	1.0151	3.90	1.0216	5.58	1.0281	7.26	1.0346	8.95
1.0022	0.57	1.0087	2.25	1.0152	3.93	1.0217	5.61	1.0282	7.29	1.0347	8.97
1.0023	0.50	1.0088	2.27	1.0153	3.95	1.0218	5.64	1.0283	7.32	1.0348	9.00
1.0024	0.62	1.0089	2.30	1.0154	3.98	1.0219	5.66	1.0284	7 - 34	1.0349	9.03
1.0025	0.64	1.0000	2.32	1.0155	4.00	1.0220	5.69	1.0285	7.37	1.0350	9.05
1.0026	0.67	1,0001	2.35	1.0156	4.03	1 0220	E 71	1.0286	7.39	1.0351	9.08
1.0027	0.69	1.0092	2.38	1.0157	4.06	1.0222	5.74	1.0287	7.42	1.0352	g.10
1.0028	0.72	1.0093	2.40	1.0158	4.08	1.0223	5 - 77	1.0288	7 - 45	1.0353	9.13
1.0029	0.75	1.0094	2.43	1.0159	4.11	1.0224	5.79	1.0289	7 - 47	1.0354	9.16
1.0030	0.77	1.0095	2 45	1.0160	4.13	1.0225	5.82				9.18
1.0031	0.80	1.0006	2.45	1.0161	4.16	1.0226	5.84	1.0290	7.50	1.0355	9.10
1.0032	0.82	1.0097	2.50	1.0162	4.19	1.0227	5.87	1.0292	7.55	1.0357	9.23
1.0033	0.85	1.0008	2.53	1.0163	4.21	1.0228	5.89	1.0293	7.58	1.0358	9.26
1.0034	0.87	1.0099	2.56	1.0164	4.24	1.0229	5.92	1.0294	7.60	1.0359	9.29
			0	6-			.				
1.0035	0.90	1.0100	2.58 2.61	1.0165	4.26	1.0230	5.94	1.0295	7.63	1.0360	9.31
1.0037	0.95	1.0102	2.63	1.0167	4.31	1.0232	5.97 6.00	1.0296	7.65	1.0361	9 · 34 9 · 36
1.0038	0.08	1.0103	2.66	1.0168	4.34	1.0233	6.02	1.0298	7.70	1.0363	9.39
1.0039	1.00	1.0104	2.69	1.0169	4.37	1.0234	6.05	1.0299	7.73	1.0364	9.42
				11			,		_	_	
1.0040	1.03	1.0105	2.71	1.0170	4.39	1.0235	6.07	1.0300	7.76	1.0365	9.44
1.0041	1.08	1.0107	2.74	1.0171	4.42	1.0230	6.10	1.0301	7.78 7.81	1.0366	9 - 47
1.0043	1.11	1.0108	2.79	1.0173	4.47	1.0238	6.15	1.0303	7.83	1.0368	9.52
1.0044	1.13	1.0109	2.82	1.0174	4.50	1.0239	6.18	1.0304	7.80	1 0309	9.55
				li	۱ ا					1	
1.0045	1.16	1.0110	2.84	1.0175	4.52	1.0240	6.20	1.0305	7.89	1.0370	9.57
1.0040	1.18	1.0111	2.89	1.0176	4.55	1.0241	6.23	1.0306	7.91	1.0371	9.60
1.0047	1.24	1.0113	2.92	1.0177	4.57	1.0243	6.25	1.0307	7 · 94 7 · 97	1.0372	9.65
1.0049	1.26	1.0114	2.94	1.0179	4.63	1.0244	6.31	1.0300	7.99	1.0374	9.68
1							-	l i		1	
1.0050	1.29	1.0115	2.97	1.0180	4.65	1.0245	6.33	1.0310	8.02	1.0375	9.70
1.0051	1.32	1.0110	3.00	1.0181	4.68	1.0246	6.36	1.0311	8.04	1.0376	9.73
1.0052	1.34	1.0117	3.02	1.0182	4.70	1.0247	6.38 6.41	1.0312	8.07	1.0377	9.75
1.0054	1.37	1.0110	3.03	1.0184	4.73	1.0240	6.44	1.0313	8.12	1.0378	9.78
			"-",	1				1		3/9	-
1.0055	1.42	1.0120	3.10	1.0185	4.78	1.0250	6.46	1.0315	8.14	1.0380	9.83
1.0056	1.45	1.0121	3.12	1.0186	4.81	1.0251	6.49	1.0316	8.17	1.0381	9.86
1.0057	1.47	1.0122	3.15	1.0187	4.83	1.0252	6.51	1.0317	8.20	1.0382	9.88
1.0058	1.50	1.0123	3.18	8810.1	4.86	1.0253	6.54	1.0318	8.22	1.0383	9.91
1.0039	1.52	1.0124	3.20	1.0189	4.00	1.0254	6.56	1.0319	8.25	1.0384	9.93
1.0060	1.55	1.0125	3.23	1.0190	4.91	1.0255	6.59	1.0320	8.27	1.0385	9.96
1.0061	1.57	1.0126	3.26	1.0191	4.94	1.0256	0.02	1.0321	8.30	1.0386	9.99
1.0062	1.60	1.0127	3.28	1.0192	4.96	1.0257	6.64	1.0322	8.33	1.0387	10.01
1.0063	1.63	1.0128	3.31	1.0193	4.99 5.01	1.0258	0.07	1.0323	8.35 8.38	1.0388	10.04
		1.0120	3.33	1.0194	3.01	1 4.0250 1	0.70	1.0324	0.38	1.0389	10.00
1.0064	05			i) .				"		1	

EXTRACT IN WINE—(Continued).

Specific Gravity.	Ex- tract.	Specific Gravity.	Ex- tract.	Specific Gravity.	Ex- tract.	Specific Gravity.	Ex- tract.	Specific Gravity.	Ex- tract.	Specific Gravity.	Ex- tract.
1.0390	10.09	1.0455	11.78	1.0520	13.47	1.0585	15.16	1.0650	16.86	1.0715	18.56
1.0391	10.11		11.81	1.0521	13.49	1.0586	15.19	1.0651	16.88	1.0716	18.58
1.0302	10.14	1.0457	11.83	1.0522	13.52	1.0587	15.22	1.0652	16.91 16.94	1.0717	18.61 18.63
1.0393	10.19	1.0459	11.88	1.0524	13.57	1.0589	15.27	1.0654	16.96	1.0719	18.66
		1				ł		1			
1.0395	10.22		11.91	1.0525	13.60	1.0590		1.0655	16.99	1.0720	18.60
1.0396	10.27	1.0461	11.00	1.0526	13.62	1.0592	15.32	1.0657	17.01	1.0721	18.71
1.0398	10.30	1.0463	11.99	1.0528	13.68	1.0503	15.37	1.0658	17.07	1.0723	18.76
1.0399	10.32	1.0464	12.01	1.0529	13.70	1.0594	15.40	1.0659	17.09	1.0724	18.79
1.0400	10.35	1.0465	12.04	1.0530	13.73	1.0595	15.42	1.0660	17.12	1.0725	18.82
1.0401	10.37	1.0466	12.06	1.0531	13.75	1.0596	15.45	1.0661	17.14	1.0726	18.84
1.0402		1.0467	12.09	1.0532	13.78	1.0597	15.48	1.0662	17.17	1.0727	18.87
1.0404	10.45		12.14	1.0534	13.83	1.0599	15.53	1.0664	17.22		18.92
								664			1
1.0405	10.48	1.0470	12.17	1.0535	13.86	1.0600	15.55	1.0665	17.25	1.0730	18.95
1.0407	10.53	1.0472	12.22	1.0537	13.91	1.0602	15.01	1.0607	17.30	1.0732	19.00
1.0408	10.56	1.0473	12.25		13.94	1.0603	15.63	1.0668	17.33	1.0733	19.03
1.0409	10.58	1.0474	12.27	1.0539	13.96	1.0604	15.66	1.0669	17.35	1.0734	19.05
1.0410	10.61	1.0475	12.30	1.0540	13.99	1.0605	15.68	1.0670	17.38	1.0735	19.08
	10.63	1.0476	12.32	1.0541	14.01	1.0606	15.71	1.0671	17.41	1.0736	19.10
1.0412	10.66	1.0477	12.35	1.0542	14.04	6-9	15.74	1.0672	17.43	1.0737	19.13
1.0414	10.71	1.0477 1.0478 1.0479	12.40	1.0544	14.09	1.0609	15.79	1.0674	17.48		19.18
1.0415	10.74	1.0480	12.43	1.0545	14.12	1.0610	15.81	1.0675	17.51	1.0740	10.21
1.0416	10.76	1.0481	12.45	1.0540	14.14	1.0611	15.84	1.0676	17.54	1.0741	19.23
1.0417	10.79	1' I.O462	12.48	1.0547	14.17	1.0612	15.87	1.0677	17.56	1.0742	19.26
1.0418	10.82	1.0483	12.51	1.0548	14.20	1.0613	15.02	1.0678	17.59	1.0743	19.29
		1.	1	1	1	1		1	1		19.31
1.0420	10.87	1.0485	12.56		14.25	1.0615	15.94	1.0680	17.64	1.0745	19.34
1.0421	10.02	1.0487		1.0552	14.30	1.0617	16.00	1.0082	17.60	1.0746	19.37
1.0423	10.95	1.0488	12.64	1.0553	14.33	1.0618	16.02	1.0683	17.72	1.0748	19.42
1.0424	10.97	1.0489	12.66	1.0554	14.35	1.0619	16.05	1.0684	17.75	1.0749	19.44
1.0425	11.00	1.0490	12.69	1.0555	14.38	1.0620		1.0685	17.77	1.0750	19.47
1.0426	11.03	1.0491	12.71		14.41	1.0621	16.10		17.80	1.0751	19.50
1.0427	11.08	1.0492	12.74	1.0557	14.43	1.0622	16.13	1.0687	17.83	1.0752	19.52
1.0429			12.79	1.0559	14.48	1.0024			17.88	1.0754	19.58
1.0430	11.13	1.0495	12.82	1.0560	14.51	1.0625	16.21	1.0600	17.00	1.0755	
1.0431		1.0496	12.84	1.0561	14.54	1.0020		1.0691	17.93	1.0756	19.60
1.0432	11.18	1.0497	12.87	1.0502	14.50	1.0027	16.20	1.0602	17.95	1.0757	19.65
I.0433 I.0434	11.21	1.0498	12.90	1.0563	14.50	1.0628	16.28		17.98	1.0758	19.68
1.0434	ì	H					_	.1	ı		19.71
1.0435	11.26	1.0500	12.95	1.0565		1.0030	16.33	1.0605	18.03	1.0760	19.73
1.0430	11.28	1.0501	12.95	1.0505	14.67	1.0031	16.30	1.0696	18.06	1.0761	19.76
1.0438	11.34	1.0503	13.03	1.0508	14.72	1.0033	16.41	1.0698	18.11	1.0763	19.79
1.0439	11.36	1.0504	13.05	1.0509	14.74	1.0634	16.44	1.0699	18.14	1.0764	19.84
1.0440			13.08	1.0570	14.77		16.47	1.0700	18.16	1.0765	19.86
1.0441	11.42	1.0500	13.10	1.0571	14.80	1.0636	16.49	1.0701	18.19	1.0706	19.89
1.0442	11.44	1.0507	13.13	1.0572	14.85	1.0638	16.52 16.54	1.0702	18.22	1.0767	19.92
1.0444		1.0509	13.18	1.0574	14.87		16.57	1.0704	18.27	1.0769	19.94
1.0445	111.52	1.0510	13.21	1.0575	14.90	1.0640	16.60	1.0705	18.30	1.0770	20.00
1.0445	11.55	1.0511	13.23	1.0576	14.93	1.0041	16.62	1.0706	18.32	1.0771	20.02
1.0447	1 44.57	1.0512	13.26	1.0577			16.65	1.0707	18.35	1.0772	20.05
1.0448	11.60	1.0513	13.29	1.0578	14.95	1.0643	16.68	1.0708	18.37	1.0773	20.07
	1			1	1	_			l i		•
1.0450	11.65	1.0515	13.34	1.0580	15.03	1.0645	16.73	1.0710	18.43	1.0775	20.12
1.0451	11.68	1.0516	13.36	1.0581	15.08	1.0647	16.75	1.0711	18.45 18.48	1.0776	20.15
1.0453	11.73	1.0518	13.42	1.0583	15.11	1.0648	16.80	1.0713	18.50	1.0778	20.20
1.0454	11.75	1.0519	13.44	1.0584	15.14	1.0649	16.83	1,0714	18.53	1.0779	20.23
~		11		Ľ'				<u> </u>	<u> </u>		

EXTRACT IN WINE—(Continued).

Specific Gravity.	Ex- tract.	Specific Gravity.		Specific Gravity.		Specific Gravity.		Specific Gravity.	Ex- tract.	Specific Gravity.	Ex- tract.
1.0780	20.26	1.0845	21.96			1.0975	25.38	1.1040	27.09	1.1105	28.8t
1.0781	20.28	1.0846	21.99	1.0011		1.0976	25.41	1.1041	27.12	1.1106	28.83
1.0782	20.31	1.0847	22.02	1.0913			25.43	1.1042	27.15	1.1107	28.86
1.0783	20.36	1.0840	22.07	1.0014		1.0078	25.46	1.1043	27.17	1.1108	28.88 28.91
		1	!			il		I'	ı	i	
1.0785		1.0850	22.00	1.0015	23.80	1.0980	25.51	1.1045	27.22	1.1110	28.94
1.0786	20.41	1.0851	22.12	1.0916	23.83	1.0981	25.54	1.1040	27.25	I.IIII	28.Q0
		1.0853	22.17	I T. COLX	23.88	1.0983	25.56 25.59	1.1047	0 20	1.1112	28.99
1.0789	20.47	1.0854	22.20	1.0019		1.0984	25.62	1.1049	27.33	1.1114	29.04
1.0790	20.52	1.0855	22.22	1.0920	23.03	1.0985	25.64	1.1050	27.35	1.1115	29.07
1.0/91		1.0856	22.25	1.0021	23.06	1.0986	25.67	1.1051	27.38	1.1116	29.09
1.0792	20.57	1.0857	22.28	1.0922	23.99	1.0987	25.70	1.1052	27.41	1.1117	29.12
1.0793	20.62	1.0858	22.30	1.0023		1.0088	25.72	1.1053		1.1118	29.15
1.0/94		Ι.	1 6	1.0924	24.04	1.0989	25.75	1.1054	27.40	1.1119	29.17
1.0795	20.65	1.0860	22.36	1.0925	24.07	1.0000	25.78 25.80	1.1055	27.49	1.1120	29.20
1.0797	20.70	1.0862	22.41	1.0927	24.00	1.0991	25.83	1.1056	27.51	1.1121	29.23 29.25
1.0798	20.73	1.0863	22.43	1.0028	24.14	1.0993	25.85	1.1058	27.57	1.1123	29.28
1.0799	20.75	1.0804	22.40	1.0929		1.0994	25.88	1.1059	27.59	1.1124	29.31
1.0800	20.78	1.0865	22.49	1.0030	24.20	1.0005	25.01	1.1060	27.62	1.1125	29.33
1.0801	20.81		22.51	1.0931	24.22	1.0996	25.93	1.1061	27.65	1.1126	29.30
1.0802	20.83	1.0867	22.54	1.0032	24.25	1.0007	25.96	1.1002	27.67	1.1127	29.39
1.0804		1.0869	22.57	1.0033	24.27	1.0000	25.99	1.1063	27.70	1.1128	29.41 29.44
1.0805	20.07	1.0870	22.62	1.0935	1	1.1000	26.04	1'			
1.0806	20.94	1.0871	22.65	1.0936	24.33 24.35		26.04 26.06		27.78	1.1130	29.47 29.49
1.0807	20.06	1.0872	22.67	1.0037	24.38	1.1002	26.00	1.1067	27.80	1.1132	20.52
1.0808	20.99	1.0873	22.70	1.0038	24.41	1.1003	20.12	1.1068	27.83	1.1133	29.54
1.0809	21.02	1.0874	22.72	1.0939	24.43	1.1004	26.14	1.1069	27.86	1.1134	29.57
1.0810		1.0875	22.75	1.0040	24.46	1.1005	26.17	1 1.1070	27.88	1.1135	29.60
1.0811	21.07	1.0876	22.78	1.0041	24.49 24.51	1.1006	26.20	1.1071	27.96	1.1136	29.62
1.0813	21.12	1.0878	22.83	1.0042	24.54	1.1007	26.25	I.1072	27.93	1.1137	29.65 29.68
1.0814	21.15	1.0870	22.80	1.0944	24.57	1.1000	26.27	1.1074	27.99	1.1139	29.70
1.0815	21.17	1.0880	22.88	1.0045	24.59	1.1010	26.30	1.1075	28.01	1.1140	20.73
1.0816	21,20	; 1.0881	22.91	1.0946	24.62	1.1011	26.33	1.1076	28.04	1.1141	29.76
1.0817	21.23	1.0882		1.0947	24.64	1.1012	26.35 26.38	1.1077	28.07	1.1142	29.78
1.0818	21.25	1.0883	22.90	1.0048	24.67	1.1013	26.41	1.1078	28.12	1.1143	29.81
1.0820		1.0885			i		!	1 1			
1.0821	21.31	1.0886	23.01	1.0050	24.72 24.75	1.1015	26.43	1.1080	28.15	1.1145	29.86 29.89
1.0822	21.36	1.0887	23.07	1.0052	24.78	1.1017	26.40	1.1082	28.20	1.1147	20.01
1.0823	21.38	1.0888			24.80	8101.1	26.51	1.1083	28 22	T TT48	29.94
1.0824	21.41	1.0889		1.0954	24.83	1.1019	26.54	1.1084	28.25	1.1149	29.96
1.0825	21.44	1.0890	23.14	1.0055	24.85	1.1020	26.56	1.1085	28.28	1.1150	29.99
1.0826	21.46	1.0890	23.17	1.0050	24.88	1.1021	26.50	1.1086	28. 30 I	1.1151	30.02
1.0827	21.49	1.0892	23.20	1.0057	24.91	1.1022	26.62	1.1087	28.33	1.1152	30.04
1.0828		1.0893	23.22	1.0958	24.93 24.96	1.1023	26.64	1.1088	28.36 28.38	1.1153	30.07 30.10
		. 1	1	1		i	1		11		
1.0830 1.0831	21 50	T 0806 :		1.0060	24.99 25.01	1.1025	26.70 26.72	1.1000	28.41	1.1155	30.13
1.0832	21.62	1.0807		1.0062	25.04	1.1027	26.75	1.1091	28.46	1.1157	30.18
1.0833	21.65	8080.1	23.35	1.0963		1.1028	26.78	1.1003	28.49	1.1158	30.21
1.0834	21.67	1.0899	23.38	1.0964	25.00	1.1029	26.80	1.1094	28.51	1.1159	30.23
1.0835	21.70	1.0000	23.41	1.0965	25.12	1.1030	26.83	1.1095	28.54	1	
1.0836	21.73	1.0901	23.43	1.0066		1.1031		1.1096	28.57	ì	
1.0837	21.75		23.46	1.0967	25.17	1.1032	20.88	1.1097	28.59	1	
1.0838	21.80	1.0904	23.49	1.0969	25.20 25.22	1.1033	26.93	1.1098	28.65		
T 0840	27 82	1.0905		1.0970	25.25	1.1035	26.96	1.1100	28.67		
1.0841	21.86	1.0906	23.54	1.0971	25.28	1.1036	26.99	1.1101	28.70		
1.0842	21.00	1.0907	23.59	1.0972	25.30	1.1037	27.01	1.1102	28.73]	
1.0843		1.0908	23.02		25.33		27.04	1.1103	28.75		
1.0844	21.94	1.0900	23.05	1.0974	25.30	1.1039	27.07	1.1104	28.78	1	

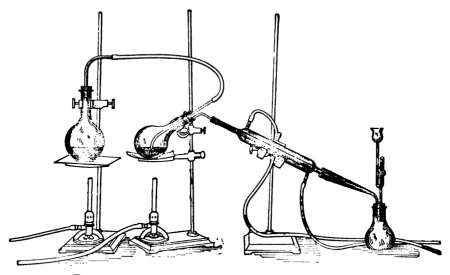


Fig. 114.—Apparatus for Determining Volatile Acids in Wine.

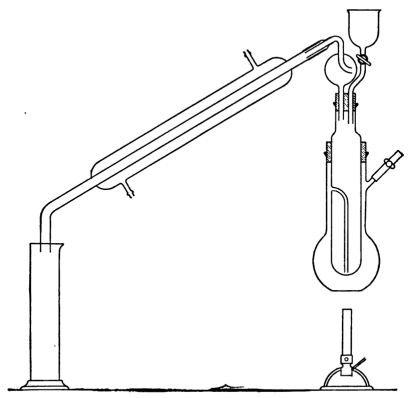


Fig. 115.—Hortvet's Apparatus for Determining the Volatile Acids in Wine.

as an indicator. Each cubic centimeter of tenth-normal alkali is equivalent to 0.006 gram acetic acid.

Hortvet Method.*—The apparatus (Fig. 115) consists of a 300 cc. flask into the neck of which is fitted a 200-cc, cylindrical flask, with a steam tube, a bulb-trap leading to a condenser, and a stop-cock funnel. The procedure is as follows: Pour 150 cc. of recently boiled water into the larger flask, attach the smaller flask by means of a section of rubber tubing, run in 10 cc. of wine (previously freed from carbonic acid). close the stop-cock and boil. In extreme cases add to the wine a small piece of paraffin to prevent foaming. When the water has boiled a moment, close the tube at the side of the larger flask and distil until 70 cc. of distillate have passed over. Transfer to a beaker, without discontinuing the distillation, and titrate, using phenolphthalein as indicator. Continue the distillation until the last 10 cc. portion requires not more than one drop of tenth-normal alkali for neutralization. Usually 80 or 90 cc. of distillate includes practically all of the volatile acids. Cool the apparatus, thus allowing the wine residue to be drawn back into the lower flask, rinse with boiled water, and reserve the total liquid for determination of non-volatile acids.

Non-volatile Acids.—These may be determined by difference, calculating the volatile acids for purposes of subtraction in terms of tartaric or other acid in which the total acidity is expressed. Non-volatile acid may be directly determined by evaporating to dryness a measured portion of the liquor, boiling the residue with water, and titrating the solution with the standard alkali.

Detection of Free Tartaric Acid.—Nessler's Method.—Some powdered cream of tartar is added to a portion of the wine in a corked flask, which is shaken from time to time, and the liquid finally filtered. To the filtrate is added a little 20% potassium acetate solution. If free tartaric acid is present, on stirring and especially after standing for some time, there will be a precipitate of cream of tartar.

Determination of Tartaric Acid, Total, Free, and Combined.—Provisional methods A. O. A. C.†

Total Tartaric Acid.—To 100 cc. of wine add 2 cc. of glacial acetic acid, 3 drops of a 20% solution of potassium acetate, and 15 grams of powdered potassium chloride, and stir to hasten solution. Add 15 cc. of 95% alcohol (specific gravity 0.81) and rub the side of the beaker vigorously with a glass rod for about one minute to start crystallization.

^{*} Jour. Ind. Eng. Chem., 1, 1909, p. 31

[†] U. S. Dept. of Agric., Bur. of Chem., Bul. 65, p. 87.

Let stand at least fifteen hours at room temperature; decant the liquid from the separated acid potassium tartrate as rapidly as possible (using vacuum) through a Gooch crucible prepared with a very thin film of asbestos, transferring no more of the precipitate to the crucible than necessary. Wash the precipitate and filter three times with a small amount of a mixture of 15 grams potassium chloride, 20 cc. of 95% alco hol (specific gravity 0.81), and 100 cc. water, using not more than 20 cc. of the wash solution in all. Transfer the asbestos film and precipitate to the beaker in which the precipitation took place, wash out the Gooch crucible with hot water, add about 50 cc. of hot water, heat to boiling, and titrate the hot solution with decinormal sodium hydroxide, using delicate litmus tincture or litmus paper as indicator. Increase the number of cubic centimeters of decinormal alkali employed by 1.5 on account of the solubility of the precipitate. This figure multiplied by 0.015 gives the amount of total tartaric acid in grams per 100 cc.

Cream of Tartar.—Ignite the residue obtained from the evaporation of 50 cc. of wine as directed under the determination of ash. Exhaust the ash with hot water, add to the filtrate 25 cc. of decinormal hydrochloric acid, heat to incipient boiling, and titrate with decinormal alkali solution, using litmus as indicator. Deduct from 25 cc. the number of cubic centimeters of decinormal alkali employed, and multiply the remainder by 0.0188 for potassium bitartrate expressed in grams.

Free Tartaric Acid.—Add 25 cc. of decinormal hydrochloric acid to the ash of 50 cc. of wine, heat to incipient boiling, and titrate with decinormal sodium hydroxide, using litmus as indicator. Deduct the number of cubic centimeters of alkali employed from 25, and multiply the remainder by 0.0075 to obtain the amount of tartaric acid necessary to combine with all the ash (considering it to consist entirely of potash). Deduct the figure so obtained from the total tartaric acid for the free tartaric acid.

Determination of Free and Combined Malic Acid in Cider and Wine.

Evaporate 100 cc. of the sample on the water-bath to half its volume, cool, and treat first with 10 cc. of 10% calcium chloride solution, and then with ammonia to strong alkaline reaction. Let stand for an hour and filter. This removes the tartaric acid. Concentrate the filtrate by evaporation on the water-bath to 25 cc., add 75 cc. of 95% alcohol, heat to the boiling-point, and filter. Wash the residue with a mixture of 3 parts 95% alcohol and 1 part water, dry, and burn to an ash. Add 25 cc. of tenth-normal hydrochloric acid to the ash, dilute with water,

heat to boiling, and titrate with tenth-normal sodium hydroxide, using phenolphthalein as an indicator. Multiply the difference between 25 and the number of cubic centimeters required to neutralize by 0.0067 for the grams of malic acid.

Polarization.—Results are usually expressed in terms of the undiluted product. The simplest method consists in treating a measured amount of wine or cider with one-tenth of its volume of lead subacetate, filtering, and polarizing the filtrate in the 200-mm. tube. The reading is increased by 10% for the true direct polarization. If the filtrate is deeply colored after clarification with subacetate, as in the case of some artificially colored wines, use a 100-mm. tube and multiply by 2 the reading, increased by 10%.

If the reducing sugars are also to be determined, one can use the same solution for both the polarization and the reducing sugars, as follows:

Exactly neutralize with sodium hydroxide solution a measured quantity of the wine, using litmus paper as an indicator, and evaporate on the water-bath to about one-fourth its original volume. Wash with water, into the measuring-glass, add enough subacetate of lead solution to clarify, and make up with water to the original volume. Filter, and to a measured amount of the filtrate add 10% of its volume of a saturated solution of sodium sulphate. Again filter and submit to polarization a portion of the filtrate, making the 10% correction on the reading.

If the invert reading is desired, subject a portion of the filtrate to inversion as described under Sugars.

Determination of Reducing Sugar.—Determine the reducing sugar as dextrose by the Defren or the Allihn method. For this purpose dilute a portion of the wine, dealcoholized and clarified as described in the preceding section, so that it contains about one-half of 1% of sugar for the Defren and about 1% of sugar for the Allihn method. One may assume 2% as the sugar-free extract of wine, the number of volumes of water to be added to the filtrate being determined by the difference between 2 and the total extract as determined.

Determination of Glycerin.—Evaporate 100 cc. of the wine on the water-bath to a volume of about 10 cc., after which 1 or 2 grams of fine sand are added and sufficient milk of lime to render alkaline. Continue the evaporation nearly to dryness, and after cooling shake with 50 cc. of 95% alcohol, heat to boiling on the water-bath, and filter. Wash the insoluble residue on the filter-paper with several portions of hot alcohol, using say 100 cc., adding the washings to the original filtrate. Evaporate the alcoholic filtrate to a syrupy consistency on the water-bath, dissolve

the residue in 10 cc. of absolute alcohol, and transfer to a flask with 15 cc. of ether. Tightly cork the flask, shake, and allow to stand for some time. Then filter into a tared dish, wash with a mixture of absolute alcohol (1 part) and ether (1.5 parts), evaporate the filtrate to syrupy consistency on the water-bath, dry in a water-oven for one hour, and weigh as glycerin.

With plastered wines the results are too high, for the reason that the potassium sulphate present is decomposed by the lime to form potassium hydroxide, soluble in glycerin and alcohol. In such wines the above residue should be ignited, and the ash deducted from the first weight.

Determination of Potassium Sulphate.—Acidify 100 cc. of the sample with hydrochloric acid, heat to boiling, and add an excess of barium chloride solution. Filter, wash, dry, ignite, and weigh as barium sulphate, calculating the equivalent of potassium sulphate. The presence of the latter in excess of 0.06 gram per 100 cc. indicates plastering.

Determination of Tannin.—An approximate method of determining tannin is that of Nessler and Barth. 12 cc. of wine are treated with 30 cc. of alcohol and filtered. 35 cc. of the filtrate, which corresponds to 10 cc. of the wine, is evaporated to about 6 cc. and transferred to a 10-cc. graduated centrifuge tube. A few drops of 40% sodium acetate are then added, and a slight excess of 10% ferric chloride. The tube is corked, gently agitated, and allowed to stand for twenty-four hours. The volume of the precipitate is then measured, each cubic centimeter being equivalent to 0.033% of tannin in the wine.

Foreign Coloring Matters in Wine.—A wide variety of artificial colors have been found in red wine. Those most commonly employed at present are cochineal, fuchsin, and acid fuchsin.

The Pharmacopæia prescribes the following color tests:

If 2 cc. of red wine be mixed in a test-tube with 2 drops of chloroform and 4 cc. of normal potassium hydroxide, and the mixture carefully heated, the disagreeable odor of isonitril should not become preceptible (absence of various anilin colors).

If 50 cc. of red wine be treated with a slight excess of ammonia water, the liquid should acquire a green or brownish-green color; if it be then well shaken with 25 cc. of ether, the greater portion of the ethereal layer removed and evaporated in a porcelain capsule with an excess of acetic acid and a few fibers of uncolored silk, the latter should not acquire a crimson or violet color (absence of fuchsin).

If 25 cc. of red wine heated to about 45° C. be well agitated with 25 grams of manganese dioxide, the liquid filtered off and acidulated with

hydrochloric acid, it should not acquire a red color (absence of sulphofuchsin).

Dupre's Method of Detection.*—Small cubes of jelly measuring about 2 cm. in thickness are prepared as follows: Dissolve 1 part of pure gelatin in 10 parts boiling water and pour upon a plate to harden. This is then cut into cubes of the above size by a sharp knife. When a wine is suspected of containing foreign color, one of the cubes is immersed therein and allowed to remain for twenty-four hours, after which it is removed, washed slightly in cold water, and cut through with a knife. If the color is a natural one, it will lightly tinge the outer surface of the cube, but will not permeate far below the surface, so that the inner portion of the cross-section will be largely free from color. Nearly all foreign coloring matters used in wine, such as most coal-tar dves, cochineal, Brazil wood, logwood, etc., will be found to deeply permeate the jelly cube often to the center. Information as to the nature of the color may sometimes be gained by immersing the dyed jelly cube in weak ammonia. If the color be rosanilin, the cube is decolorized, if cochineal, a purple coloration will result, and if logwood, a brown tinge.

Cazeneuve's Methods for Detecting Colors in Wine.—While by no means complete, the following method of Cazeneuve as condensed and arranged by Gautier (La Sophistication des Vins) will often be found helpful. If other colors than these are evidently present, tests should be made as indicated in Chapter XVII. Cazeneuve employs the following reagents:

- (1) Yellow oxide of mercury, finely pulverized.
- (2) Lead hydrate, freshly precipitated, well washed, suspended in about twice its volume of water; to be kept in a stoppered bottle; should be renewed after several days' use.
- (3) Gelatinous ferric hydrate, well washed from ammonia, suspended in about twice its volume of water.
 - (4) Manganese dioxide, pulverized.
 - (5) Concentrated, chemically pure sulphuric acid.
 - (6) White wool.
- (7) Stannous hydrate, freshly precipitated, well washed, suspended in water, and kept from exposure to light and air.
- (8) Collodion silk, the artificial silk produced from nitro-cellulose. This fiber has a special affinity for basic dyes.

^{*} Jour. Chem. Soc., 37, p. 572.

To 10 cc. of the wine are added 0.2 gram finely powdered yellow oxide of mercury. Boil and pour upon a double filter.

E	Filtrate	colored vello	W. 10 CC.	Filtrate colored		Filtrate colorless after acidifying.
Filtrate colored blue. boiled with collodion s	Filtrate co	lored yellow.	ilter.	cc. of the wine are with 2 grams lead and filtered.	l hydrate	Ten cc. treated with iron hydrate to boiling.
The wine	hydrate the liqui	is added and d is boiled.	rate is color which aft with pure ed is	The preceding soluted monia. Acidify and Wash the fibers, dry, concentrated H ₂ SO ₄ , lows:	The preceding solution decolorized by ammonia.	of wine are ro grams of and brought
Fiber is colored blue	solution colored yealow, but only intly. Wool fibers are dyed, ashed, dried, and treated with the concentrated H ₂ SO ₄ . Color oduced is	Solution colorless. Wool fibers are dyed, washed, dried, and treated with pure concentrated H ₂ SO ₄ . Color produced is	red red. Boil up with wool er washing and drying are concentrated H ₂ SO,. Color	The preceding solution is not changed by ammonia. Acidify and boil up with wool fibers. Wash the fibers, dry, and treat them with pure concentrated H ₂ SO ₄ , which colors them as follows:	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Filtrate colorless { i
Ammonia is ad from its natural c mercury. Causes cipitate.	Brown-yellow	Brownyellow	Fuchsin red	:	To a portion of the wine is added its weight of manganese dioxide. Acidify and filter.	Ten cc. wine are treated with 20 grams stannous hydrate and brought to boiling. A fluorescent rose
nnia is added to the wine freed natural color by yellow oxide of . Causes a blue to yellow pre-		on diluting			yellowish. Filtrate yellow-red or red.	s hy- boil- Colored
Methylene blus	. Yellow NS . Marlius yellow	Chrysoin Veswin lluting Solid yellow Yellow N	Orange I and a Orange I and a Tropeolin o Tropeolin y. Tropeolin oo (Orange IV) Helianthin, Orange III	Roccellin Pur ple-red Bordeaux red Ponceaux Biebrich scarlet Croceine 3B Croceine 7B	Ruchsin Acid Juchsin	Pure wine Wine colored with voge- table pigmonts Cockineal Estin Erythrosin

MALT LIQUORS. BEER.

In its widest sense beer may be defined as the product of fermentation of an infusion of almost any farinaceous grain with various bitter extractives, but unless otherwise qualified it should be strictly applied to the beverage resulting from the fermentation of malted barley and hops. In the manufacture of beer two distinct processes are employed, viz., malting or sprouting the grain, and brewing. Many brewers do nothing but the latter, buying their malt already prepared.

Malting.—For the preparation of malt, the barley is steeped in water for several days, after which the water is drained off and the moist grain is "couched," or piled in heaps, on a cement floor, where it undergoes a spontaneous heating process, during which it germinates, forming the ferment diastase. When the maximum amount of diastase has been produced, indicated by the length of growth of the sprout, or "acrospire" within the grain, the germination is checked by spreading the grain in layers over a perforated iron floor, and finally subjecting it to artificial heat. The character of the malt and of the beer produced from it depends largely on the heat at which the "green" malt is kiln dried. If dried between 32° and 37° C. it forms pale malt, which produces the lightest grades of beer. Most beer is made from malt dried at higher temperatures, say from 38° to 50°, the depth of color of the liquor varying with the heat to which the malt has been subjected, while the color of the malt varies from the "pale" through the "amber" to "brown," or even black. The darkest grades are sometimes dried at temperatures over 100° C., even to the point where the starch becomes caramelized.

A more modern method consists in the so-called pneumatic malting, wherein the whole operation is conducted in a large rotating drum, which holds the grain, and in which the temperature and moisture at different stages is carefully controlled by the admission to the interior of the drum of moisture-laden or dry air, heated to the required degree.

The chief object of malting is the production of diastase, which by its subsequent action on the starch converts it into the fermentable sugars maltose and dextrin. Malt contains much more diastase than is necessary to convert the starch simply contained therein to maltose, and is capable of acting on the starch of a considerable quantity of raw grain, such as corn or rice, when mixed with it. This practice of using other grains than malt is prohibited in some localities, such as Bavaria.

Brewing.—The malt, or mixture of malt and raw grain, is first crushed and "mashed" by stirring with water in tubs at 50° to 60° C., finally heating to 70°. During this process the conversion of the starch to maltose and dextrin takes place. The resulting liquor is known as "wort." containing, besides maltose and dextrin, peptones and amides. clear wort is then drawn off from the residue, and boiled to concentrate the product and to sterilize it, after which hops (the female flower of the Humulus lubulus) are added and the boiling continued. contain resins, bitter principles, tannic acid, and a peculiar essential oil, all of which are to some extent imparted to the wort. After cooling and settling, the clear wort is run into fermenting-vats, where selected veast, usually saccharomyces cerevisiae, is added, and the alcoholic fermentation allowed to proceed. The temperature greatly affects the character of the fermentation. If kept between 5° and 8° C., a slow fermentation proceeds, known as bottom fermentation, during which the veast settles out at the bottom. This is much more easily controlled than the quick or top fermentation, which takes place at from 15° to 18°. much of the yeast in the latter case being carried to the surface, from which it is finally removed by skimming. In either case the yeast feeds upon the albuminous matter present. At the proper stage the beer is drawn off from the larger portion of the yeast, and run into casks, or tuns, in which an after-fermentation proceeds. The beer is finally clarified by treatment with gelatin or beech shavings or chips, to which the floating yeast-cells and other impurities attach themselves. It is finally stored in barrels coated with brewers' pitch, or pasteurized at 60° C. and bottled.

Varieties of Beer.—Formerly the division of beers into "lager," "schenk," and "bock" was made by reason of the fact that beer had to be brewed under certain climatic conditions and at certain seasons only. Now, with improved means for artificial refrigeration, and with better methods controlling all stages of the process, these distinctions are less marked.

Lager Beer (from lager, a storehouse) is a term originally applied to Bavarian beer, but is now given to any beer that has been stored several months. Formerly lager beer was made early in the winter, and stored in cool cellars till the following spring or summer, during nearly all of which time a slow after-fermentation took place. The best lager beers contain a low proportion of hops, and are high in extract and alcohol.

Schenk Beer is a quickly fermented beer made in winter for immedi-

ate use. It is brewed in from four to six weeks and will not keep long without souring.

Bock Beer, according to older systems of nomenclature, occupied a middle place between lager and schenk, being an extra strong beer brewed for spring use and made in limited quantities, not being intended for storage.

Berlin Weiss Bier is prepared by the quick or top fermentation of a wort consisting of a mixture of malted barley and wheat with hops. It is high in carbon dioxide, being usually bottled before the second fermentation has ended.

Ale is virtually the English name for beer. It is usually lighter colored than lager beer, being made from pale malt by quick or top fermentation, and containing rather more hops than beer. It has a high content of sugar, due to checking fermentation at an earlier stage than in ordinary beer.

Porter is a dark ale, the deep color of which should be due to the use of brown malt dried at a high temperature, but which is sometimes colored by the admixture of caramel. It has a large extract, chiefly sugar.

Stout is an extra-strong porter, being high both in alcohol and extract. Composition of Beer.—Beer is a somewhat complex liquor. Besides water, alcohol, and sugar, it contains carbon dioxide, succinic acid, dextrin, glycerin, tannic acid, the resinous bitter principles of hops, nitrogenous bodies (chiefly peptones and amides), alkaline and lime salts (chiefly phosphates), fat (traces), acetic acid and lactic acid. The latter acid constitutes the chief fixed acid of beer.

The following analyses of different varieties of beer are due to König:

Variety.	Number of Analyses.	Specific Gravity.	Water.	Carbonic Acid.	Alcohol by Weight.	Extract.	Nitrogenous Substances.	Sugar as Maltose.	Gum and Dextrin.	Acid as Lactic.	Glycerin.	Ash.	Phosphoric Acid.
Schenk Lager Export beer. Bock Weiss bier Porter	258 109 84 26 40	1.0114 1.0162 1.0176 1.0213 1.0137 1.0191	90.08 89.01 87.87 91.63 88.49	0.196 0.209 0.234 0.297 0.215	3.93 4.40 4.69 2.73 4.70	5.79 6.38 7.21 5.34 6.59	0.71 0.74 0.73 0.58 0.65	0.88 1.20 1.81 1.62 2.62	3.73 3.47 3.97 2.42 3.08	0.151 0.161 0.165 0.392	0.165 0.154 0.176 0.092	0.247 0.263 0.149 0.363	0.07 7 0.074 0.089

Fifteen samples of lager beer and seven samples of pale ale, bought in Massachusetts bar-rooms, representing as nearly as possible the quality

of liquor sold	every day t	o patrons by	the bottle or	glass, were	analyzed
by the Board	of Health w	ith the follow	ing results:		-

		Per Cent of Original Wort Extract.	Per Cent of Alcohol by Weight.	Per Cent of Extract.
Beer—	Maximum	18.91	7.07	7.76
	Minimum.	7-33	1.10	7.76 3.67
	Mean	15.04	4 - 45	5.92
Pale ale	—Maximum	15.99	5-37	5.47
	Minimum.	10.95	3-53	5 · 47 3 · 38 4 · 54
	Mean	13.56	4.49	4.54

Five out of the 15 beer samples and 3 out of the 7 ale samples contained salicylic acid.

The percentage composition of the ash of German beer is thus given by König as the mean of 10 analyses:

Ash in 100 Parts Beer.	Potash.	Soda.	Lime.	Magnesia.	Iron Oxide.	Phos- phoric Acid.	Sul- phuric Acid.	Silica.	Chlorine,
0.306	33.67	8.94	2.78	6.24	0.48	31.35	3 · 47	9.29	2.93

Malt and Hop Substitutes.—By reason of the fluctuation in market price of these two chief constituents of beer, it sometimes becomes a question of economy to employ cheaper substitutes wholly or in part for one or the other. There are two classes of malt substitutes, (1) those which, like corn, rice, and wheat, are mixed directly with the malt before "mashing," and, like the malt, have to undergo a saccharous fermentation before being acted on by yeast, and (2) such substances as cane sugar, invert sugar, commercial glucose, and dextrin, which are added to the wort at a later stage in the brewing, just before the addition of the yeast, being in condition to be readily acted on by the latter.

Glucose is by far the most common malt substitute, by reason of the fact that its sugars much resemble those of malt, and are in readily fermentable form. Diastase forms from the malt dextrin and maltose, while commercial glucose contains dextrin, maltose, and dextrose.

When the price of malt is abnormally high, the addition of glucose is decidedly economical, but when ordinary conditions prevail, the cost of the two, figured with reference to their yield in alcohol and extract, is about the same. Aside from the question of economy, however, there are advantages in the use of glucose, such as diminishing the nitrogenous content of the wort without lessening the alcohol or extract yielded.

The nitrogenous matter left after fermentation is one of the chief causes of cloudiness or turbidity in the finished product, and is sometimes difficult to remove. By the use of glucose, especially in brewing clear bottled ales and sparkling pale beers, the appearance of the product is much enhanced. The temptation at times to add more glucose than is necessary to accomplish this is great. A high-grade malt may have as much as 40% of glucose added to its wort and still produce an acceptable beer. With a low-grade malt, glucose yields a very poor quality of beer. Hence the use of glucose may or may not be desirable, though it can hardly be considered unqualifiedly as an adulterant.

As to the employment of hop substitutes, the question of relative price again enters in. Only when the price of hops is high is there any special inducement to use substitutes. Quassia wood, chiretta, gentian, and calumba, all of which yield bitter principles, have been used in beer, and cannot be considered detrimental to health. Allen and Chattaway have found the first two in beer examined by them.* Such poisonous substances as cocculus indicus, picric acid, and strychnine are alleged to have been used as hop substitutes, but there is no authentic record of any of them having been found in recent years, if at all.

Adulteration of Malt Liquors and Standards of Purity.—The Joint Committee on Standards of the A. O. A. C. and the A. S. N. F. D. D. has adopted the following standards:

Malt Liquor is a beverage made by the alcoholic fermentation of an infusion, in potable water, of barley malt and hops, with or without unmalted grains or decorticated and degerminated grains.

Beer is a malt liquor produced by bottom fermentation, and contains in 100 cc., at 20° C., not less than 5 grams of extractive matter and 0.16 gram of ash, chiefly potassium phosphate, and not less than 2.25 grams of alcohol.

Lager Beer, Stored Beer, is beer which has been stored in casks for a period of at least three months, and contains, in 100 cc., at 20° C., not less than 5 grams of extractive matters, and 0.16 gram of ash, chiefly potassium phosphate, and not less than 2.50 grams of alcohol.

Malted Beer is beer made of an infusion, in potable water, of barley, malt, and hops, and contains, in 100 cc., at 20° C., not less than 5 grams of extractive matter, nor less than 0.2 gram of ash, chiefly potassium phosphate, not less than 2.25 grams of alcohol, nor less than 0.4 gram of crude protein (nitrogen × 6.25).

^{*} Analyst, 12, 112.

• Ale is a malt liquor produced by top fermentation, and contains, in 100 cc., at 20° C., not less than 2.75 grams of alcohol, nor less than 5 grams of extract, and not less than 0.16 gram of ash, chiefly potassium phosphate.

Porter and Stout are varieties of malt liquors made in part from highly roasted malt.

Non-injurious bitter principles are no doubt employed in place of hops, and unless the liquor is sold for a pure malt beer, they cannot be regarded as adulterants.

The tendency to shorten the time of storage of beer, or to sell it without storing at all, lessens or does away with the after-fermentation, resulting in a lack of "life" or effervescence in the product. This is sometimes made up by the addition of sodium bicarbonate.

Distinction between Malted and Non-malted Liquors.—In some states where strict prohibitory liquor laws are in force, it is illegal to sell "malt liquors," so that when convictions are obtained, it is necessary for the analyst to distinguish between liquors brewed wholly or in part from malt and those in which no malt has been used, but which were brewed entirely from malt substitutes. This distinction is not always easy to make with precision. In the absence of malt, glucose is usually the sole source of alcohol in these beverages. Parsons * has shown that the most striking points of difference between malted and non-malted liquors are in their per cent of phosphoric acid and albuminoids, and that pure malt beer or ale should contain at least 0.04% P₂O₆, and 0.25% albuminoids (N×6.25). A low ash and high content of sulphates in the ash are also indicative of glucose. The following analyses made by Parsons clearly show these distinctions:

COMPOSITION OF SEVENTY-SIX SAMPLES OF AMERICAN MALT LIQUORS.

	Specific Gravity.	Alcohol by Vol- ume.	Extract.	Albumin- oids (N×6.25)	Phos- phoric Acid.	Ash.	Sul- phates in Ash.	Free Acid.
Average Maximum Minimum		5.61 7.85 0.35	4.61 7.64 3.15	0.470 0.614 0.290	0.061 0.095 0.045	0.209 0.296 0.147	6.34 12.67 2.44	0.26 0.87 0.10

^{*} Jour. Am. Chem. Soc., 24, 1902, p. 1179.

TYPICAL ANALYSES OF BEERS APPARENTLY NOT BREWED FROM MALT.

Number.	Specific Gravity.	Alcohol by Vol- ume.	Extract.	Albumin- oids (N×6.25)	Phos- phoric Acid.	Ash.	Sul- phates.	Pree Acid.
I	1.0074	1.68 2.63	2.52 3.40	0.114	0.010	0.19	21.22	Normal
3	1.00Ó2 1.0112	2.27 2.11	2.25 3.53	0.150	0.015	0.124 0.140	11.30	**
5	1.0041	1.85	1.73	0.031	0.010	0.088	12.50	**

The ash of the fifth sample is thus compared with that of the average beer as given by Blyth:

• • •	Malt Beer (Blyth).	"No-malt" Beer (Parsons).
K ₂ O	37.22	12.93
Na ₂ O	8.04	19.61
CaO	1.93	Undetermined
MgO	5.51	"
Fe ₂ O ₃	Trace	"
SO ₃	1.44	10.81
P ₂ O ₈	32.09	10.71
Cl	2.91	21.76
SiO ₂	10.82	7-50

Preservatives in Beer.—Antiseptics are frequently added to malt liquors, salicylic acid being most commonly used. Fluorides of ammonium and sodium have been found in American beer. Other preservatives to be looked for are benzoic acid and sulphites. Beer casks are frequently "sulphured" or fumed with a solution of calcium bisulphite, so that the beer may derive its content of sulphites from this source.

In cases of police seizure of beer sold in bulk or in opened bottles for the purpose of ascertaining whether or not their alcoholic content exceeds certain limits fixed by law, a little formalin had best be added as soon as possible after the seizure to prevent further fermentation. This is especially desirable in cases where there is likely to be some delay in making the analysis, so as to forestall any claim on the part of the defendant of additional alcohol being formed after the seizure. From 6 to 8 drops of a 40% solution of formaldehyde to a quart of beer is sufficient, and this quantity will not appreciably affect the analysis.

Arsenic in Beer.—In 1900 a very disastrous epidemic of arsenical poisoning occurred in Manchester, England, involving several thousand cases, many of which were fatal. The arsenic was traced to sulphuric

acid, which entered into the manufacture of commercial glucose used in the beer, the acid found so highly arsenical being made from a certain variety of Swedish pyrites, which was found to be abnormally high in arsenic. There appeared to be no doubt whatever that the beer was the sole cause of the trouble. While the presence of arsenic was in this case accidental, carelessness was shown on the part of those having to do with the purity of the materials entering into the composition of the beer. Fortunately no other instances are on record of arsenical poisoning from malted liquors. A large number of samples of American beer have been examined in the laboratory of the Food and Drug Department of the Massachusetts State Board of Health, and only insignificant traces of arsenic have in any case been found.

Temperance Beers and Ales.—Many varieties of these so-called temperance drinks are home-made, as well as sold on the market. They are usually slightly fermented infusions of various roots and herbs, including ginger or sassafras, with molasses or sugar and yeast, and more often contain less than 1% of alcohol by volume. Among them are included spruce beer, and the various root beers, such as ginger beer and ginger ale. The latter beverages are generally carbonated. Numerous brands of bottled beer are manufactured, which contain virtually the same body and characteristic flavor as lager beer, but not the alcohol. Indeed the composition of many of these beverages is identical with that of lager beer, excepting in alcoholic content, being made by substantially the same process and out of the same ingredients, but with the alcohol finally removed by steaming, so that the liquor comes within the limits of a temperance beverage. Of this class is *Uno beer*, which ranges from 0.6 to 0.9 per cent in alcohol.

METHODS OF ANALYSIS OF MALT LIQUORS.*

• Preparation of Sample.—Transfer the contents of the bottle or bottles to a large flask and shake vigorously to hasten the escape of carbon dioxide, care being taken that the liquor is not below 15° C., since below this temperature the carbon dioxide is retained by the beer and is liable to form bubbles in the pycnometer.

Specific Gravity.—See page 657.

Ash.—Determine in 25 cc. by evaporation and ignition at dull redness.

^{*} Barnard, U. S. Dept. of Agric., Bur. of Chem., Circ. 33. A. O. A. C. Methods, ibid., Bul. 107 (rev.), p. 90.

Determination of Alcohol.—From the Specific Gravity of the Distillate.—Proceed as described on p. 658, employing 100 cc. of the liquor, and determining the specific gravity at 15.5° C. If the liquor is markedly acid, add 0.1 to 0.2 gram of precipitated calcium carbonate previous to distillation.

From the Refraction of the Distillate.—Prepare the distillate as described on p. 658, except that it is made up to the mark at 17:5° C. Determine the refraction at 17.5° C. by means of the immersion refractometer, and calculate the alcohol by the table of Ackermann and Steinmann below.

ACKERMANN AND STEINMANN'S TABLE FOR OBTAINING THE PER-CENTAGE OF ALCOHOL IN THE DISTILLATE OF BEER FROM THE IMMERSION REFRACTOMETER READINGS.*

						1		1	1	
Refractometer Reading. Alcohol by Weight,	Alcohol by Volume, Per Cent.	Refractometer Reading.	Alcohol by Weight, Per Cent.	Alcohol by Volume, Per Cent.	Refractometer Reading.	Alcohol by Vight, Per Cent.	Alcohol by Volume, Per Cent.	Refractorneter Leading.	Alcohol by Voight, Per Feat.	Alcohol by Volume, Per Cent.
15.0 0.00 15.1 0.00 15.2 0.13 15.3 0.19 15.4 0.21 15.5 0.38 15.7 0.44 15.8 0.50 15.9 0.50 16.1 0.70 16.2 0.77 16.3 0.83 16.4 0.88 16.5 0.90 16.7 1.00 16.7 1.00 16.8 1.10 17.0 1.22 17.1 1.33	0.08 0.16 0.24 0.40 0.48 0.56 0.64 0.72 0.80 0.88 0.96 1.04 1.12 1.19 1.27 1.33 1.43 1.51 1.58	17.2 17.3 17.4 17.5 17.6 17.7 17.8 17.9 18.0 18.1 18.3 18.4 18.5 18.6 18.7 18.8 18.9 19.0	1.38 1.44 1.51 1.57 1.63 1.68 1.74 1.81 1.87 1.93 2.06 2.13 2.19 2.25 2.31 2.37 2.43 2.49 2.55 2.68	1.74 1.82 1.90 1.98 2.05 2.12 2.20 2.28 2.36 2.44 2.52 2.60 2.68 2.76 2.84 2.99 3.07 3.14 3.29 3.37	19.4 19.5 19.6 19.7 19.8 19.9 20.0 20.1 20.2 20.3 20.4 20.5 20.6 20.7 20.8 20.9 21.0 21.1 21.2 21.3 21.5	0.730 0.00 0.00 0.00 0.00 0.00 0.00 0.00	0.455168 1030 0.7744 100 0.71774844 4.444	21.6 21.0 01.0 01.0 00.0 00.3 20.4 22.5 20.7 20.0 20.0 20.1 20.0 20.1 20.0 20.1 20.0 20.1 20.0 20.1 20.0 20.0	4.02 4.07 4.13 4.22 4.33 4.39 4.44 4.54 4.54 4.70 4.81 4.81 4.92 4.97 5.02	5.06 5.13 5.20 5.26 5.32 5.32 5.53 5.55 5.78 5.78 5.78 5.78 6.00 6.07 6.20 6.27 6.33

† Zeits. gesamte Brauwesen, 28, 1905, p. 259.

Determination of Extract.—In cases where extreme accuracy is desired, the result obtained by evaporating at 100° a weighed amount of the beer cannot be accepted, on account of the dehydration of the maltose at a temperature exceeding 75° C. Unless the evaporation is conducted at that temperature (a difficult operation), a closer approxi-

EXTRACT IN BEER WORT.* [According to Schultz and Ostermann,]

	Ext	ract.		Ext	ract.		Ext	ract.		Ext	ract.
Specific Gravity at 15° C.	Per Cent by Weight	Grams per 100 cc.	Specific Gravity at 15° C.	Per Cent by Weight	Grams per 100 cc.	Specific Gravity at 15° C.	Per Cent by Weight	Grams per 100 cc.	Specific Gravity at 15° C.	Per Cent by Weight	Grams per 100 cc.
1.0000	0.00	0.00	1.0065	1.60	1.70	1.0130	3 - 35	3 - 39	1.0195	5.06	5.16
1.0001,	0.03	0.03	1.0006	1.72	1.73	1.0131	3.38	3.42	1.0196	5.09	5.19
1.0002	0.05	0.05	1.0067	1.74	1.75	1.0132	3.41	3.46	1.0197	5.12	5.22
1.0003	0.10	0.10	1.0069	1.77	1.80	1.0134	3.43 3.46	3.48 3.51	1.0199	5.15	5.25
1.0005	0.13	0.13	1.0070	1.82	1.83	1.0135	3.48	3.53	1.0200	5.20	.5.30
1.0006	0.16	0.16	1.0071	1.84	1.85	1.0136	3.51 3.54	3.56 3.59	1.0201	5.23	5.34 5.36
1.0008	0.21	0.21	1.0073	1.90	1.01	1.0138	3.56	3.61	1.0203	5.28	5.39
1.0009	0.24	0.24	1.0074	1.92	1.93	1.0139	3 · 59	3.64	1.0204	5.30	5.41
1.0010	0.26	0.26	1.0075	1.95	1.96	1.0140	3.61	3.66	1.0205	5.33	5.44
1.0011	0.29	0.29	1.0076	1.97	2.02	1.0141	3.64 3.66	3.69	1.0206	5.35 5.38	5.46
1.0012	0.31	0.31	1.0077	2.00	2.04	1.0143	3.69	3.71 3.74	1.0208	5.40	5.49 5.51
1.0014	0.37	0.37	1.0079	2.05	2.07	1.0144	3.72	3.77	1.0209	5 · 43	5.54
1.0015	0.39	0.39	1.0080	2.07	2.09	1.0145	3.74	3.79 3.83	1.0210	5.45 5.48	5.56 5.60
1.0010	0.42	0.42	1.0081	2.10	2.14	1.0147	3·77 3·79	3.85	1.0211	5.50	5.62
1.0018	0.47	0.47	1.0083	2.15	2.17	1.0148	3.82	3.88	1.0213	5.53	5.65
1.0019	0.50	0.50	1.0084	2.17	2.19	1.0149	3.85	3.91	1.0214	5.55	5.67
1.0020	0.52	0.52	1.0085	2.20	2.22	1.0150	3.87	3.93	1.0215	5 . 57	5.69
1.0021	0.55	0.55	1.0086	2.23	2.25	1.0151	3.90	3.96 3.98	1.0216	5.60	5.72 5.74
1.0022	0.60	0.58	1.0088	2.25	2.30	1.0153	3.95	4.01	1.0218	5.65	5.77
1.0024	0.63	0.63	1.0089	2.30	2.32	1.0154	3.97	4.03	1.0219	5.67	5.79
1.0025	0.66	0.66	1.0090	2.33	2.35	1.0155	4.00	4.06	1.0220	5.70	5.83
1.0026	0.68	0.68	1,0001	2.35	2.37	1.0156	4.03	4.09	1.0221	5.72	5.85 5.88
1.0027	0.71	0.71	1.0092	2.38	2.40	1.0157	4.05	4.11	1.0222	5 · 75 5 · 77	5.90
1.0029	0.76	0.76	1.0094	2.43	2.45	1.0159	4.10	4.17	1.0224	5.80	5.93
1.0030	0.79	0.79	1.0095	2.46	2.48	1.0160	4.13	4.20	1.0225	5.82 5.84	5.95 5.97
1.0031 1.0032	0.84	0.84	1.0090	2.48	2.50	1.0162	4.18	4.25	1.0227	5.87	6.00
1.0033	0.87	0.87	1.0098	2.53	2.55	1.0163	4.21	4.28	1.0228	5.89	6.02
1.0034	0.89	0.89	1.0099	2.56	2.59	1.0164	4.23	4.30	1.0229	5.92	6.06
1.0035	0.92	0.92	1.0100	2.58	2.61	1.0165	4.26	4.33	1.0230	5.94	6.08
1.0036 1.0037	0.94	0.94	1.0101	2.61	2.64	1.0166	4.28	4.35	1.0231	5.97 5.99	6.11
1.0038	1.00	1.00	1.0103	2.66	2.60	1.0168	4.31 4.34	4.41	1.0233	6.02	6.16
1.0039	1.02	1.02	1.0104	2.69	2.72	1.0169	4.36	4.43	1.0234	6.04	6.18
I.0040 I.004I	1.05	1.05	1.0105	2.71	2.74	1.0170	4.39	4.46	1.0235	6.07 6.09	6.21 6.23
1.0042	1.10	1.10	1.0107	2.76	2.70	1.0172	4.44	4.52	1.0237	6.11	6.25
1.0043	1.13	1.13	8010.1	2.79	2.79	1.0173	4.47	4.55	1.0238	6.14	6.29
1.0044	1.15	1.16	1.0109	2.82	2.85	1.0174	4.50	4.58	1.0239	6.16	6.31
1.0045	1.18	1.19	1.0110	2.84	2.87	1.0175	4.53	4.61	1.0240	6.19	6.34
1.0046 1.0047	1.21	I.22 I.24	1.0111	2.87	2.90	1.0176	4.55 4.58	4.63	1.0241	6.24	6.39
1.0048	1.26	1.27	1.0113	2.92	2.95	1.0178	4.61	4.69	1.0243	6.26	6.41
1.0049	1.29	1.30	1.0114	2.94	2.97	1.0179	4.63	4.71	1.0244	6.29	6.44
1.0050	1.31	1.32	1.0115	2.97	3.00	1.0180	4.66	4-74	1.0245	5.31	6.46
1.0051	1.34	1.35	1.0116	3.02	3.02	1.0181	4.69 4.71	4.77	1.0246	6.34	6.50
1.0053	1.30	1.40	1.0118	3.05	3.00	1.0183	4.74	4.83	1.0247	6.39	6.55
1.0054	1.41	1.42	1.0119	3.07	3.11	1.0184	4.77	4.86	1,0249	6.41	6.57
1.0055 1.0056	1.44	1.45	1.0120	3.10	3.14 3.16	1.0185	4.79	4.88	1.0250	6.44	6.60 6.63
1.0050	1.46	I.47 I.50	1.0121	3.12	3.10	1.0180	4.82	4.Ç. 4.94	1.0251	6.47	6.66
1.0058	1.51	1.52	1.0123	3.17	3.21	1.0188	4.88	4.97	1.0253	6.52	6.68
1.0059	1.54	1.55	1.0124	3.20	3.24	1.0189	4.90	4.99	1.0254	6.55	6.72
1.0060	1.56	1.57	1.0125	3.23	3.27	1.0190	4.93	5.02	1.0255	6.58	6.75
1.0061 1.0062	1.59	1.60	1.0126	3.25	3.29	1.0191	4.98	5.05 5.08	1.0256	6.61 6.63	6.78 6.80
2.0063	1.64	1.65	1.0128	3.30	3.34	1.0193	5.01	5.11	1.0258	6.66	6.83
1.0064	1.67	1.68	1.0129	3.33	3.37	1.0194	5.04	5.14	1.0259	6.69	6.86
	·	<u> </u>	loulated f	'		<u>'</u>	<u> </u>			<u> </u>	

^{*} Calculated from results obtained by drying below 75° C.

EXTRACT IN BEER WORT-(Continued).

	Ext	ract.		Ext	ract.		Ext	ract.		Ext	ract.
Specific Gravity at 15° C.	Per Cent by Weight	Grams per 100 cc.	Specific Gravity at 15° C.	Per Cent by Weight	Grams per 100 cc.	Specific Gravity at 15° C.	Per Cent by Weight	Grams per 100 cc.	Specific Gravity at 15° C.	Per Cent by Weight	Grams per 100 cc.
I.0260 I.0261 I.0262 I.0263 I.0264	6.71 6.74 6.77 6.80	6.88 6.92 6.95 6.98	1.0325 1.0326 1.0327 1.0328	8.27 8.29 8.32 8.34	8.54 8.56 8.59 8.61	1.0390 1.0391 1.0392 1.0393	9.92 9.95 9.97 9.99	10.34 10.36 10.38	1.0455 1.0456 1.0457 1.0458	11.53 11.55 11.57 11.60	12.05 12.08 12.10 12.13
1.0265 1.0266 1.0267 1.0268	6.82 6.85 6.88 6.91 6.93	7.00 7.03 7.00 7.09 7.12	1.0329 1.0330 1.0331 1.0332 1.0333	8.40 8.43 8.45 8.48	8.65 8.68 8.71 8.73 8.76	1.0394 1.0395 1.0396 1.0397 1.0398	10.04 10.06 10.09 10.11	10.41 10.44 10.46 10.49 10.51	1.0459 1.0460 1.0461 1.0462 1.0463	11.62 11.65 11.67 11.70 11.72	12.15 12.19 12.21 12.24 12.26
1.0269 1.0270 1.0271 1.0272	6.96 6.99 7.01 7.04	7.15 7.18 7.20 7.23	1.0334 1.0335 1.0336 1.0337	8.51 8.53 8.56 8.59	8.82 8.85 8.88	1.0399 1.0400 1.0401 1.0402	10.13 10.16 10.18 10.20	10.53 10.57 10.59 10.61	1.0464 1.0465 1.0466 1.0467	11.75 11.77 11.79 11.82	12.30 12.32 12.34 12.37
I.0273 I.0274 I.0275 I.0276 I.0277	7.07 7.10 7.12 7.15 7.18	7.26 7.29 7.32 7.35 7.38	1.0338 1.0339 1.0340 1.0341 1.0342	8.61 8.64 8.67 8.70 8.72	8.90 8.93 8.96 9.00 9.02	1.0403 1.0404 1.0405 1.0406 1.0407	10.23 10.25 10.27 10.30 10.32	10.64 10.66 10.69 10.72	1.0468 1.0469 1.0470 1.0471 1.0472	11.84 11.87 11.89 11.92 11.94	12.39 12.43 12.45 12.48 12.50
1.0278 1.0279 1.0280 1.0281	7.21 7.23 7.26 7.28	7.41 7.43 7.46 7.48	1.0343 1.0344 1.0345 1.0346	8.75 8.78 8.80 8.83	9.05 9.08 9.10 9.14	1.0408 1.0409 1.0410 1.0411	10.35 10.37 10.40 10.42	10.77 10.79 10.83 10.85	I.0473 I.0474 I.0475 I.0476	11.97 11.99 12.01 12.04	12.54 12.56 12.58 12.61
1.0282 1.0283 1.0284	7.30 7.33 7.35	7.51 7.54 7.56 7.58	1.0347 1.0348 1.0349	8.86 8.88 8.91	9.17 9.19 9.22 9.25	1.0412 1.0413 1.0414	10.45 10.47 10.50	10.88 10.90 10.93	1.0477 1.0478 1.0479 1.0480	12.06	12.64 12.67 12.69
1.0286 1.0287 1.0288 1.0289	7.39 7.42 7.44 7.46	7.60 7.63 7.65 7.68	1.0351 1.0352 1.0353 1.0354	8.97 8.99 9.02 9.05	9.28 9.31 9.34 9.37	1.0416 1.0417 1.0418 1.0419	10.55 10.57 10.60 10.62	10.99 11.01 11.04 11.06	1.0481 1.0482 1.0483 1.0484	12.16 12.19 12.21 12.23	12.74 12.78 12.80 12.82
1.0291 1.0292 1.0293 1.0294	7.48 7.51 7.53 7.55 7.57	7.70 7.73 7.75 7.77 7.79	1.0355 1.0356 1.0357 1.0358 1.0359	9.07 9.10 9.13 9.15 9.18	9.39 9.42 9.46 9.48 9.51	1.0420 1.0421 1.0422 1.0423 1.0424	10.65 10.67 10.70 10.72 10.75		1.0485 1.0486 1.0487 1.0488 1.0489	12.28 12.31 12.33 12.36	12.85 12.88 12.91 12.93 12.96
1.0295 1.0296 1.0297 1.0298 1.0299	7.60 7.62 7.64 7.66 7.69	7.82 7.85 7.87 7.89 7.92	1.0360 1.0361 1.0362 1.0363 1.0364	9.21 9.24 9.26 9.29 9.31	9.54 9.57 9.60 9.63 9.65	1.0425 1.0426 1.0427 1.0428 1.0429	10.77 10.80 10.82 10.85 10.88	11.23 11.26 11.28 11.31 11.35	1.0490 1.0491 1.0492 1.0493 1.0494	12.38 12.41 12.43 12.45 12.48	12.99 13.02 13.04 13.06 13.10
I.0300 I.0301 I.0302 I.0303 I.0304	7.71 7.73 7.75 7.77 7.80	7.94 7.96 7.98 8.01 8.04	1.0365 1.0366 1.0367 1.0368 1.0369	9.34 9.36 9.38 9.41 9.43	9.68 9.70 9.72 9.76 9.78	1.0430 1.0431 1.0432 1.0433 1.0434	10.90 10.93 10.95 10.98 11.00	11.37 11.40 11.42 11.46 11.48	1.0495 1.0496 1.0497 1.0498 1.0499	12.50 12.53 12.55 12.58 12.60	13.12 13.15 13.17 13.21 13.23
1.0305 1.0306 1.0307 1.0308 1.0309	7.82 7.84 7.86 7.89 7.91	8.06 8.06 8.10 8.13 8.15	1.0370 1.0371 1.0372 1.0373 1.0374	9.45 9.48 9.50 9.52 9.55	9.80 9.83 9.85 9.88 9.91	1.0435 1.0436 1.0437 1.0438 1.0439	11.03 11.05 11.08 11.10 11.13		1.0500 1.0501 1.0502 1.0503 1.0504	12.63 12.66 12.67 12.70 12.72	13.26 13.28 13.31 13.34 13.36
1.0310 1.0311 1.0312 1.0313 1.0314	7.93 7.95 7.98 8.00 8.02	8.18 8.20 8.23 8.25 8.27	1.0375 1.0376 1.0377 1.0378 1.0379	9.57 9.59 9.62 9.64 9.66	9.93 9.95 9.98 10.00	I.0440 I.0441 I.0444 I.0443 I.0444		11.64 11.67 11.70 11.73 11.75	1.0505 1.0506 1.0507 1.0508 1.0509	12.75 12.77 12.80 12.82 12.85	13.39 13.42 13.45 13.47 13.50
1.0315 1.0316 1.0317 1.0318 1.0319	8.04 8.07 8.09 8.11 8.13	8.29 8.33 8.35 8.37 8.39	1.0380 1.0381 1.0382 1.0383 1.0384	9.69 9.71 9.73 9.76 9.78	10.06 10.08 10.10 10.13 10.16	1.0445 1.0446 1.0447 1.0448 1.0449	11.28 11.30 11.33 11.35 11.38	11.78 11.80 11.84 11.86 11.89	1.0510 1.0511 1.0512 1.0513 1.0514	12.87 12.90 12.92 12.94 12.97	13.53 13.56 13.58 13.60 13.64
1.0320 1.0321 1.0322 1.0323 1.0324	8.16 8.18 8.20 8.22 8.23	8.42 8.44 8.46 8.49 8.52	1.0385 1.0386 1.0387 1.0388 1.0389	9.81 9.83 9.85 9.88 9.90	10.19 10.21 10.23 10.20	1.0450 1.0451 1.0452 1.0453 1.0454	11.48	11.91 11.95 11.97 12.00	1.0515 1.0516 1.0517 1.0518 1.0519	12.99 13.02 13.04 13.07 13.09	13.66 13.69 13.71 13.75 13.77

EXTRACT IN BEER WORT-(Continued).

	Ext	ract.		Ext	ract.		Bxt	ract.		Ext	ract.
Specific Gravity at 15° C.	Per Cent by Weight	Grams per 100 cc.	Specific Gravity at 15° C.	Per Cent by Weight	Grams per 100 cc.	Specific Gravity at 15° C.	Per Cent by Weight	Grams per 100 cc.	Specific Gravity at 15° C.	Per Cent by Weight	Grams per 100 cc.
1.0520	13.12	13.80	1.0585	14.75	15.61	1.0650	16.25	17.31	1.0715	17.81	19.08
1.0521	13.14	13.82	1.0586	14.78	15.65	1.0651	16.27	17.33	1.0716	17.84	19.12
1.0522	13.16	13.85	1.0587	14.81	15.68	1.0652	16.30	17.36	1.0717	17.86	19.14
1.0523	13.19	13.88	1.0588	14.83	15.70	1.0653	16.32	17.39	1.0718	17.88	19.16
1.0524	13.21	13.90	1.0589	14.85	15.74	1.0654	16.35	17.42	1.0719	17.90	19.19
1.0525	13.24	13.94	1.0590	14.89	15.77	1.0655	16.37	17.44	1.0720	17.93	19.22
1.0526	13.26	13.96	1.0591	14.91	15.79	1.0656	16.40	17.48	1.0721	17.95	19.24
1.0527	13.29	13.99	1.0592	14.94	15.82	1.0657	16.42	17.50	1.0722	17.97	19.27
1.0528	13.31	14.01	1.0593	14.96	15.85	1.0658	16.45	17.53	1.0723	17.99	19.29
1.0529	13.34	14.05	1.0594	14.99	15.88	1.0659	16.47	17.56	1.0724	18.02	19.32
1.0530 1.0531 1.0532 1.0533 1.0534	13.36 13.38 13.41 13.43 13.46	14.07 14.09 14.12 14.15 14.18	1.0595 1.0596 1.0597 1.0598 1.0599	15.02 15.04 15.07 15.09 15.11	15.91 15.94 15.97 15.99 16.02	1.0660 1.0661 1.0662 1.0663 1.0664	16.50 16.52 16.54 16.57 16.59	17.59 17.61 17.63 17.67 17.69	1.0725 1.0726 1.0727 1.0728 1.0729	18.04 18.06 18.08 18.11 18.13	19.35 19.37 19.39 19.43
1.0535	13.48	14.20	1.0600	15.14	16.05	1.0665	16.62	17.73	1.0730	18.15	19.47
1.0536	13.51	14.23	1.0601	15.16	16.07	1.0666	16.64	17.75	1.0731	18.17	19.50
1.0537	13.53	14.26	1.0602	15.18	16.09	1.0667	16.67	17.78	1.0732	18.20	19.53
1.0538	13.56	14.29	1.0603	15.20	16.12	1.0668	16.69	17.80	1.0733	18.22	19.55
1.0539	13.58	14.31	1.0604	15.23	16.15	1.0669	16.72	17.84	1.0734	18.24	19.58
1.0540 1.0541 1.0542 1.0543 1.0544	13.61 13.63 13.66 13.68 13.71	14.34 14.37 14.40 14.42 14.46	1.0605 1.0606 1.0607 1.0608	15.25 15.27 15.29 15.31 15.34	16.17 16.20 16.22 16.24 16.27	1.0670 1.0671 1.0672 1.0673 1.0674	16.74 16.76 16.79 16.81 16.84	17.86 17.88 17.92 17.94 17.98	1.0735 1.0736 1.0737 1.0738 1.0739	18.26 18.29 18.31 18.33 18.35	19.60 19.64 19.66 19.68 19.71
1.0545	13.73	14.48	1.0610	15.36	16.30	1.0675	16.86	18.00	I.0740	18.38	19.74
1.0546	13.76	14.51	1.0611	15.38	16.32	1.0676	16.89	18.03	I.074I	18.40	19.76
1.0547	13.78	14.53	1.0612	15.40	16.34	1.0677	16.91	18.05	I.0742	18.42	19.79
1.0548	13.81	14.57	1.0613	15.43	16.38	1.0678	16.94	18.09	I.0743	18.44	19.81
1.0549	13.83	14.59	1.0614	15.45	16.40	1.0679	16.96	18.11	I.0744	18.47	19.84
1.0550	13.86	14.62	1.0615	15.47	16.42	1.0680	16.99	18.15	1.0745	18.49	19.87
1.0551	13.88	14.64	1.0616	15.49	16.44	1.0681	17.01	18.17	1.0746	18.51	19.89
1.0552	13.91	14.68	1.0617	15.52	16.48	1.0682	17.03	18.19	1.0747	18.53	19.91
1.0553	13.93	14.70	1.0618	15.54	16.50	1.0683	17.06	18.23	1.0748	18.55	19.94
1.0554	13.96	14.73	1.0619	15.56	16.52	1.0684	17.08	18.25	1.0749	18.57	19.96
1.0555 1.0556 1.0557 1.0558 1.0559	13.98 14.01 14.03 14.06 14.08	14.76 14.79 14.81 14.84 14.87	1.0620 1.0621 1.0622 1.0623 1.0624	15.58 15.60 15.63 15.65 15.67	16.55 16.57 16.60 16.62 16.64	1.0685 1.0686 1.0687 1.0688 1.0689	17.11 17.13 17.16 17.18	18.28 18.31 18.34 18.36 18.40	1.0750 1.0751 1.0752 1.0753 1.0754	18.59 18.62 18.64 18.66 18.68	
1.0560	14.11	14.90	1.0625	15.69	16.66	1.0690	17.23	18.42	1.0755	18.70	20.11
1.0561	14.13	14.92	1.0626	15.72	16.70	1.0691	17.25	18.44	1.0756	18.72	20.14
1.0562	14.16	14.96	1.0627	15.74	16.73	1.0692	17.28	18.48	1.0757	18.74	20.16
1.0563	14.18	14.98	1.0628	15.76	16.75	1.0693	17.30	18.50	1.0758	18.76	20.18
1.0564	14.21	15.01	1.0629	15.78	16.77	1.0694	17.33	18.53	1.0759	18.78	20.21
1.0565	14.23	15.03	1.0630	15.80	16.80	1.0695	17.35	18.56	1.0760	18.81	20.24
1.0566	14.26	15.07	1.0631	15.83	16.83	1.0696	17.38	18.59	1.0761	18.83	20.26
1.0567	14.28	15.09	1.0632	15.85	16.85	1.0697	17.40	18.61	1.0762	18.85	20.29
1.0568	14.31	15.12	1.0633	15.87	16.87	1.0698	17.43	18.65	1.0763	18.87	20.31
1.0569	14.33	15.15	1.0634	15.89	16.90	1.0699	17.45	18.67	1.0764	18.89	20.33
1.0570 1.0571 1.0572 1.0573 1.0574	14.36 14.38 14.41 14.44 14.46	15.18 15.20 15.23 15.27 15.29	1.0635 1.0636 1.0637 1.0638 1.0639	15.92 15.94 15.96 15.98 16.01	16.93 16.95 16.98 17.00 17.03	1.0700 1.0701 1.0702 1.0703 1.0704	17.48 17.50 17.52 17.54 17.57	18.70 18.73 18.75 18.77 18.81	1.0765 1.0766 1.0767 1.0768 1.0769	18.91 18.93 18.95 18.97	20.36 20.38 20.40 20.43 20.46
1.0575	14.49	15.32	1.0640	16.03	17.06	1.0705	17.59	18.83	1.0770	19.02	20.48
1.0576	14.52	15.36	1.0641	16.05	17.08	1.0706	17.61	18.85	1.0771	19.04	20.51
1.0577	14.54	15.38	1.0642	16.07	17.10	1.0707	17.63	18.88	1.0772	19.06	20.53
1.0578	14.57	15.41	1.0643	16.09	17.12	1.0708	17.66	18.91	1.0773	19.08	20.55
1.0579	14.59	15.43	1.0644	16.12	17.16	1.0709	17.68	18.93	1.0774	19.10	20.58
1.0580	14.62	15.47	1.0645	16.14	17.18	1.0710	17.70	18.96	1.0775	19.12	20.60
1.0581	14.65	15.50	1.0646	16.16	17.20	1.0711	17.72	18.98	1.0776	19.14	20.63
1.0584	14.67	15.52	1.0647	16.18	17.23	1.0712	17.75	19.01	1.0777	19.17	20.66
1.0583	14.70	15.56	1.0648	16.21	17.26	1.0713	17.77	19.04	2.0778	19.19	20.68
1.0584	14.73	15.59	5.0649	16.23	17.28	1.0714	17.79	19.06	1.0779	19.21	20.71

EXTRACT IN BEER WORT-(Continued).

	Ext	ract.		Ext	ract.		Ext	ract.	!	Ext	ract.
Specific Gravity at 15° C.	Per Cent by Weight	Grams per 100 cc.	Specific Gravity at 15° C.	Per Cent by Weight	Grams per 100 cc.	Specific Gravity at 15° C.	Per Cent by Weight	Grāms per 100 cc.	Specific Gravity at 13° C.	Per Cent by Weight	Grams per 100 cc.
1.0780	19.23	20.73	1.0845	20.70	22.45	1.0910	22.19	24.21	1.0975	23.59	25.89
1.0781	19.25	20.75	1.0846	20.73	22.48	1.0911	22.21	24.24	1.0976	23.61	25.92
1.0782	19.27	20.78	1.0847	20.75	22.50	1.0912	22.23	24.26	1.0977	23.63	25.94
1.0783	19.29	20.80	1.0848	20.77	22.53	1.0913	22.26	24.29	1.0978	23.65	25.97
1.0784	19.31	20.82	1.0849	20.79	22.55	1.0914	22.28	24.31	1.0979	23.67	25.99
1.0785 1.0786 1.0787 1.0788 1.0789	19.33 19.36 19.38 19.40 19.42	20.85 20.88 20.90 20.93 20.95	1.0850 1.0851 1.0852 1.0853 1.0854	20.81 20.83 20.86 20.88 20.90	22.58 22.61 22.64 22.66 22.68	1.0015 1.0016 1.0017 1.0018	22.30 22.32 22.34 22.37 22.39	24.34 24.37 24.30 24.42 24.44	1.0980 1.0981 1.0982 1.0983	23.69 23.71 23.73 23.76 23.78	26.01 26.04 26.06 26.09 26.11
1.0790	19.44	20.98	1.0855	20.93	22.72	1.0920	22.41	24.47	1.0985	23.80	26.14
1.0791	19.46	21.00	1.0856	20.95	22.75	1.0921	22.43	24.49	1.0986	23.82	26.17
1.0792	19.49	21.03	1.0857	20.98	22.78	1.0922	22.45	24.51	1.0987	23.84	26.19
1.0793	19.51	21.06	1.0858	21.01	22.81	1.0923	22.48	24.54	1.0988	23.86	26.22
1.0794	19.53	21.08	1.0859	21.04	22.84	1.0924	22.50	24.56	1.0989	23.88	26.24
1.0795 1.0796 1.0797 1.0798 1.0799	19.56 19.58 19.60 19.63 19.65	21.11 21.14 21.16 21.20 21.22	1.0860 1.0861 1.0862 1.0863 1.0864	21.06 21.09 21.11 21.13 21.16	22.87 22.90 22.93 22.96 22.99	1.0925 1.0926 1.0927 1.0928 1.0929	22.52 22.54 22.56 22.59 22.61	24.60 24.62 24.64 24.67 24.70	1.0990 1.0991 1.0992 1.0993	23.90 23.92 23.94 23.97 23.99	26.27 26.30 26.32 26.35 26.37
1.0800	19.67	21.24	1.0865	21.19	23.02	1.0930	22.63	24.73	1.0995	24.01	26.40
1.0801	19.70	21.28	1.0866	21.22	23.06	1.0931	22.65	24.76	1.0996	24.03	26.42
1.0802	19.72	21.30	1.0867	21.25	23.09	1.0932	22.67	24.78	1.0997	24.05	26.44
1.0803	19.74	21.33	1.0868	21.28	23.12	1.0933	22.69	24.81	1.0998	24.07	26.47
1.0804	19.77	21.36	1.0869	21.30	23.15	1.0934	22.71	24.83	1.0999	24.09	26.49
1.0805	19.79	21.38	1.0870	21.33	23.18	1.0935	22.73	24.86	1.1000	24.11	26.52
1.0806	19.81	21.41	1.0871	21.35	23.21	1.0936	22.75	24.89	1.1001	24.13	26.55
1.0807	19.84	21.43	1.0872	21.37	23.23	1.0937	22.77	24.91	1.1002	24.15	26.57
1.0808	19.86	21.46	1.0873	21.39	23.26	1.0938	22.80	24.93	1.1003	24.17	26.60
1.0809	19.88	21.49	1.0874	21.41	23.28	1.0939	22.82	24.96	1.1004	24.19	26.62
1.0810 1.0811 1.0812 1.0813 1.0814	19.91 19.93 19.96 19.98 20.00	21.52 21.55 21.58 21.60 21.63	1.0875 1.0876 1.0877 1.0878 1.0879	21.43 21.45 21.47 21.40 21.51	23.31 23.33 23.36 23.38 23.40	1.0940 1.0941 1.0942 1.0943	22.84 22.86 22.88 22.90 22.92	24.99 25.01 25.03 25.06 25.08	1.1005 1.1006 1.1007 1.1008 1.1009	24.21 24.23 24.25 24.28 24.30	26.65 26.68 26.70 26.73 26.75
1.0815	20.03	21.66	1.0880	21.54	23.43	1.0945	22.94	25.11	1.1010	24.32	26.78
1.0816	20.05	21.69	1.0881	21.56	23.45	1.0946	22.96	25.14	1.1011	24.34	26.81
1.0817	20.07	21.71	1.0882	21.58	23.48	1.0947	22.98	25.16	1.1012	24.36	26.83
1.0818	20.10	21.74	1.0883	21.60	23.50	1.0948	23.00	25.18	1.1013	24.30	26.86
1.0819	20.12	21.77	1.0884	21.62	23.52	1.0949	23.03	25.21	1.1014	24.41	26.88
1.0320	20.14	21.79	1.0885	21.64	23.55	1.0950	23.05	25.24	1.1015	24.43	26.91
1.0821	20.17	21.83	1.0886	21.66	23.58	1.0951	23.07	25.26	1.1016	24.45	26.93
1.0822	20.19	21.85	1.0887	21.68	23.60	1.0952	23.10	25.20	1.1017	24.47	26.95
1.0823	20.21	21.87	1.0888	21.71	23.63	1.0953	23.12	25.31	1.1018	24.49	26.98
1.0824	20.24	21.91	1.0889	21.73	23.66	1.0954	23.14	25.34	1.1019	24.51	27.00
1.0825	20.26	21.93	1.0890	21.75	23.69	1.0955	23.16	25.37	1.1020	24.53	27.03
1.0826	20.28	21.96	1.0891	21.77	23.72	1.0956	23.18	25.39	1.1021	24.55	27.06
1.0827	20.31	21.99	1.0892	21.70	23.74	1.0957	23.20	25.42	1.1022	24.57	27.08
1.0828	20.33	22.01	1.0893	21.82	23.77	1.0958	23.23	25.45	1.1023	24.60	27.11
1.0829	20.35	22.04	1.0894	21.84	23.79	1.0959	23.25	25.47	1.1024	24.62	27.14
1.0830	20.37	22.06	1.0895	21.86	23.82	1.0960	23.27	25.50	1.1025	24.64	27.17
1.0831	20.39	22.08	1.0896	21.89	23.85	1.0961	23.29	25.53	1.1026	24.66	27.19
1.0832	20.41	22.11	1.08)7	21.91	23.87	1.0962	23.31	25.55	1.1027	24.68	27.21
1.0833	20.43	22.13	1.0898	21.93	23.90	1.0963	23.33	25.58	1.1028	24.70	27.24
1.0834	20.46	22.16	1.0899	21.96	23.93	1.0964	23.35	25.60	1.1029	24.72	27.26
1.0835	20.48	22.19	1.0900	21.98	23.96	1.0965	23.37	25.63	1.1030	24.74	27.29
1.0836	20.50	22.21	1.0901	22.00	23.98	1.0966	23.39	25.66	1.1031	24.76	27.32
1.0837	20.52	22.24	1.0902	22.02	24.01	1.0967	23.41	25.68	1.1032	24.78	27.34
1.0838	20.54	22.26	1.0903	22.04	24.03	1.0968	23.44	25.71	1.1033	24.81	27.37
1.0839	20.56	22.29	1.0904	22.06	24.05	1.0969	23.46	25.73	1.1034	24.83	27.39
1.0840 1.0841 1.0842 1.0843 1.0844	20.50 20.62 20.64 20.66 20.68	22.32 22.35 22.38 22.40 22.42	1.0905 1.0906 1.0907 1.0908 1.0909	22.08 22.10 22.12 22.15 22.17	24.08 24.11 24.13 24.16 24.18	1.0970 1.0971 1.0072 1.0073	23.48 23.50 23.52 23.55 23.55	25.76 25.79 25.81 25.84 25.86	1.1035 1.1036 1.1037 1.1038 1.1039	24.85 24.87 24.89 24.92 24.94	27.42 27.45 27.47 27.50 27.53

EXTRACT IN BEER WORT-(Concluded).

	Ext	ract.		Ext	ract.		Ext	ract.		Ext	ract.
Specific Gravity at 15° C.	Per Cent by Weight	Grams per 100 cc.	Specific Gravity at 15° C.	Per Cent by Weight	Grams per 100 cc.	Specific Gravity at 15° C.	Per Cent by Weight	Grams per 100 cc.	Specific Gravity at 15° C.	Per Cent by Weight	Grams per 100 cc.
1.1040	24.96	27.56	1.1095	26.16	29.03	1.1150	27.29	30.43	1.1205	28.38	31.81
1.1041	24.98	27.58	1.1096	26.18	29.06	1.1151	27.31	30.45	1.1206	28.40	31.83
1.1042	25.00	27.60	1.1097	26.20	29.08	1.1152	27.33	30.47	1.1207	28.42	31.86
1.1043	25.03	27.63	1.1098	26.23	29.11	1.1153	27.35	30.50	1.1208	28.44	31.88
1.1044	25.05	27.66	1.1099	26.25	29.13	1.1154	27.37	30.52	1.1209	28.46	31.90
1.1045 1.1046 1.1047 1.1048	25.07 25.09 25.11 25.14 25.16	27.69 27.72 27.74 27.77 27.70	1.1100 1.1101 1.1102 1.1103 1.1104	26.27 26.29 26.31 26.33 26.35	29.16 29.19 29.21 29.24 29.26	1.1155 1.1156 1.1157 1.1158 1.1159	27.38 27.40 27.42 27.44 27.46	30.55 30.57 30.59 30.62 30.64	1.1210 1.1211 1.1212 1.1213 1.1214	28.48 28.50 28.52 28.54 28.56	31.93 31.95 31.98 32.00 32.03
1.1050	25.18	27.82	1.1105	26.37	29.29	1.1160	27.48	30.67	1.1215	28.58	32.05
1.1051	25.20	27.85	1.1106	26.39	29.32	1.1161	27.50	30.69	1.1216	28.60	32.08
1.1052	25.22	27.87	1.1107	26.41	29.34	1.1162	27.52	30.72	1.1217	28.62	32.11
1.1053	25.24	27.90	1.1108	26.44	29.37	1.1163	27.54	30.75	1.1218	28.64	32.13
1.1054	25.27	27.93	1.1100	26.46	29.39	1.1164	27.56	30.77	1.1210	28.66	32.15
1.1055	25.29	27.96	1.1110	26.48	29.42	1.1165	27.58	30.80	I.1220	28.68	32.18
1.1056	25.31	27.98	1.1111	26.50	29.44	1.1166	27.60	30.82	I.1221	28.70	32.20
1.1057	25.33	28.00	1.1112	26.52	29.46	1.1167	27.62	30.85	I.1222	28.72	32.23
1.1058	25.35	28.03	1.1113	26.54	29.49	1.1168	27.64	30.87	I.1223	28.74	32.25
1.1059	25.38	28.06	1.1114	26.56	29.51	1.1169	27.66	30.89	I.1224	28.76	32.27
1.1060 1.1061 1.1062 1.1063 1.1064	25.40 25.42 25.44 25.46 25.48	28.09 28.12 28.14 28.17 28.10	1.1115 1.1116 1.1117 1.1118 1.1110	26.58 26.60 26.62 26.64 26.66	29.54 29.57 29.59 29.61 29.64	1.1170 1.1171 1.1172 1.1173 1.1174	27.68 27.70 27.72 27.74 27.76	30.92 30.94 30.97 31.00	1.1225 1.1226 1.1227 1.1228 1.1220	28.78 28.80 28.82 28.84 28.86	32.30 32.32 32.35 32.37 32.40
1.1065 1.1066 1.1067 1.1068 1.1069	25.50 25.52 25.54 25.57	28.22 28.25 28.27 28.30 28.32	1.1120 1.1121 1.1122 1.1123 1.1124	26.68 26.70 26.72 26.75 26.77	29.67 29.69 29.71 29.74 29.77	1.1175 1.1176 1.1177 1.1178 1.1179	27.78 27.80 27.82 27.84 27.86	31.05 31.07 31.09 31.12 31.15	1.1230 1.1231 1.1232 1.1233 1.1234	28.88 28.90 28.92 28.94 28.94	32.43 32.45 32.48 32.50 32.53
1.1070 1.1071 1.1072 1.1073 1.1074	25.59 25.61 25.63 25.65 25.67 25.69	28.35 28.38 28.40 28.43 28.45	1.1125 1.1126 1.1127 1.1128 1.1120	26.79 26.81 26.83 26.85 26.85	29.80 29.83 29.85 29.88 29.90	1.1180 1.1181 1.1182 1.1183 1.1184	27.88 27.90 27.92 27.94 27.96	31.18 31.20 31.23 31.25 31.27	1.1235 1.1236 1.1237 1.1238 1.1239	28.98 29.00 29.02 29.04	32.56 32.58 32.60 32.63 32.65
1.1075	25.71	28.48	1.1130	26.89	29.93	1.1185	27.98	31.30	1.1240	29.08	32.68
1.1076	25.73	28.51	1.1131	26.91	29.95	1.1186	28.00	31.32	1.1241	29.10	32.71
1.1077	25.75	28.53	1.1132	26.93	29.97	1.1187	28.02	31.35	1.1242	29.12	32.73
1.1078	25.78	28.56	1.1133	26.95	30.00	1.1188	28.04	31.37	1.1243	29.14	32.76
1.1079	25.78	28.56	1.1134	26.97	30.02	1.1189	28.07	31.40	1.1244	29.16	32.76
1.1080	25.82	28.61	1.1135	26.99	30.06	1.1190	28.09	31.43	I.1245	20.18	32.81
1.1081	25.84	28.64	1.1136	27.01	30.08	1.1191	28.11	31.45	I.1246	29.20	32.83
1.1082	25.86	28.66	1.1137	27.03	30.10	1.1192	28.13	31.48	I.1247	29.22	32.86
1.1083	25.89	28.69	1.1138	27.05	30.13	1.1193	28.15	31.51	I.1248	29.24	32.89
1.1084	25.91	28.72	1.1139	27.07	30.15	1.1194	28.17	31.53	I.1240	20.26	32.91
1.1085	25.93	28.75	I.II40	27.09	30.18	1.1195	28.19	31.56	1.1250	29.28	32.94
1.1086	25.96	28.78	I.II41	27.11	30.20	1.1196	28.21	31.59	1.1251	29.30	32.96
1.1087	25.98	28.80	I.II42	27.13	30.22	1.1197	28.23	31.61	1.1252	29.32	32.99
1.1088	26.01	28.83	I.II43	27.15	30.25	1.1198	28.25	31.63	1.1253	29.34	33.02
1.1089	26.03	28.83	I.II44	27.17	30.27	1.1199	28.27	31.65	1.1254	29.36	33.04
1.1090	26.05	28.89	1.1145	27.10	30.31	1.1200	28.28	31.68	1.1255	29.38	33.27
1.1091	26.07	28.92	1.1146	27.21	30.33	1.1201	28.30	31.70	1.1256	29.40	33.09
1.1092	26.09	28.94	1.1147	27.23	30.35	1.1202	28.32	31.73	1.1257	29.42	33.12
1.1093	26.12	28.97	1.1148	27.25	30.37	1.1203	28.34	31.75	1.1258	29.45	33.14
1.1094	26.14	29.00	1.1149	27.27	30.40	1.1204	28.36	31.78	1.1259	29.47	33.17

ACKERMANN'S TABLE FOR OBTAINING THE EXTRACT IN BEER FROM THE IMMERSION REFRACTOMETER

	Ex- tract in 100 cc. Grams.	7.97	7.99	8.03	8.05	8.07	8.10	8.12	8.15	8.17	8.30	8.33	8.25	8.28	8.30	8.33	8.35	8.38	8.40	8.43	8.46
	R-R.	31.0	н	n	٣	4	'n	9	7	∞	6	32.0	н	"	8	4	'n	9	7	∞	6
	'Ex- tract in 100 cc. Grams.	7.45	7.48	7-51	7.53	7.56	7.58	7.61	7.63	7.66				7.76	7.79	7.81	7.84	7.87	7.89	7.92	7.94
R (R').*	R-R'.	29.0	+	8	6	4	'n	9	7	∞	6	30.0	1	7	es	4	'n	۰.	7	∞	٥.
3 BEER	Ex- tract in 100 cc. Grams.	6.94	6.97	86.0	7.01	7.03	7.06	7.08	7.11	7.14	7.16	7.20	7.22	7.25	7.27	7.30	7-33	7.35	7.38	7.40	7-43
OF THE BEER (R').*	R-R'.	27.0	H	0	٣	4	v	9	7	∞	٥	28.0	н	"	8	4	S	9	7	∞	6
	Ex- tract in roo cc. Grams.	6.43	6.45	6.48	6.50	6.53	6.55	6.58	19.9	6.63	99.9	6.68	6.71	6.73	92-9	6.79	6.8I	6.84	98.9	6.89	16.9
DISTILLATE	R-R'.	25.0	-	71	~	4	s	9	7	∞	6	26.0		а	3	4	S	9	7	∞	6
	Extract in 100 cc. Grams.	5.91	5.94	2.96	5.09	10.9	6.04	6.07	6.00	6.12	6.14	6.17	6.19	6.22	6.25	6.27	6.30	6.32	6.35	6.37	6.40
AND OF THE	R-R'	23.0	H	a	3	4	25	9	7	∞	6	24.0	H	64	3	4	S	9	7	∞	6
(R) AND	Extract in 100 cc. Grams.	5.40	5.42	5-45	5.47	5.50	5.53	5.55	5.58	5.60	5.63	5.65	5.68	5.71	5.73	5.76	5.78	5.81	5.83	5.86	5.89
EER (A	R-R'.	21.0	н	~	3	4	25	9	7	•	6	22.0	-	~	3	4	S	9	7	œ	6
I AININ THE B	Extract in 100 cc. Grams.	4.88	16.4	4.93	4.96	4.98	5.01	5 04	3.06	5.00	5.11	5.14	5.17	5.19	5.22	5.24	5.27	5.29	5.32	5.35	5-37
G OF	R-R'.	19.0	н	"	8	4	10	9	~	· ••	6	20.0	-	64	3	4	2	9	7	∞	6
EADIN	Extract in 100 cc. Grams.	4 37	4.40	4.43	4.45	4 47	4.50	4.52	4.55	4.57	9.4	4.63	4.65	4.68	4.70	4.73	4.75	4.78	18.4	4.83	4.86
S IAI	R-R'.	17.0	н	"	3	4	v	9	7	· •	6	18.0	н	61	<i>ب</i>	4	Ŋ	9	7	«	6
ACKERMANN'S IABLE FOR OBIAINING THE READING OF THE BEER (R)	Extract in 100 cc. Grams.	3.76	3.88	3.91	3.93	3.96	3.98	10.4	4.04	4.06	4.09	4.11	4.14	4.16	4.19	4.22	4.24	4.27	4.29	4.32	4.34
ACKE	RIK	15.0	н	"	3	4	ĸ	9	7	∞	6	0.01	н	a	8	4	v	9	7	œ	0,

* Zeits. gesamte Brauwesen, 29, 1906, p. 146.

mation to the truth is obtained, especially with beer high in sugar, by calculation as follows:

From the Specific Gravity.—Evaporate a measured quantity of the beer to one-fourth its volume on the water-bath, make up with water to its original measure, and determine the specific gravity of the deal-coholized beer. Then by means of Schultz and Ostermann's table, pp. 716-20, calculate the extract corresponding.

From the Refraction.—Method of Ackermann and Foggenburg.
—Determine the refraction of the liquor at 17.5° C. by means of the immersion refractometer. Determine also the refraction of the distillate from 100 cc. of the liquor at 17.5° C. after making up to its original volume. In order to secure accurate results, care should be taken to cool the prism of the instrument to exactly 17.5° C. by immersing for five minutes in the water-bath previous to taking the refraction of the liquids. If determinations are made on a number of samples, this cooling is not necessary except before taking the reading of the first of the series.

Calculate the grams of extract (E) from the refraction of the liquor (R) and of the distillate (R') by the following formula:

$$E = 0.25705(R - R')$$
.

The extract is more conveniently obtained from Ackermann's table given on p. 721.

Original Gravity of Beer Wort and its Determination.—Following a long-established custom of the English excise, the duty on beer has been based on the specific gravity of the original wort, by which is meant the wort of the beer before any of its sugar has been lost by fermentation.

From the content of alcohol in the beer the sugar originally present in the wort may be calculated, assuming that the alcohol amounts to about half the sugar used up in fermentation.

Obtain the specific gravity of the beer, dealcoholized and made up to its original volume, as in the calculation of the extract. This is called the "extract gravity." Note the specific gravity corresponding to the alcohol found, i.e., the specific gravity of the distillate in the alcohol determination, when made up to the original volume, and subtract this from 1. The difference is known arbitrarily as the "degree of spirit indication."

From the table of Graham, Hofmann, and Redwood,* p. 723, the "degrees of gravity lost" corresponding to the "spirit indication"

^{*} Report on Original Gravities, 1852; Allen's Com. Org. Anal., I., p. 136.

are ascertained. This figure is added to the "extract gravity" to find the "original gravity of the wort."

Degrees of "Spirit In- dication."	#.0000	0.0001	0.0002	0.0003	0.0004	0.0005	0.0006	0.0007	0.0008	0.0009
0.000		0.0003	0.0006	0.0009	0.0012	0.0015	0.0018	0.0021	0.0024	0.0027
.001	.0030	.0033	.0037	.0041	.0044	.0048	.0051	.0055	.0059	.0062
-002	.0066	.0070	.0074	.0078	.0082	.0086	.0000	.0094	.9098	.0102
.003	.0107	.0111	.0115	.0120	.0124	-0129	.0133	.0138	.0142	-0147
-004	.0151	.0155	.0160	.0164	.0168	.0173	.0177	.0182	.0186	.0191
-005	.0195	.0199	.0204	.0200	.0213	-0218	.0222	.0227	.0231	.0236
.006	.0241	-0245	.0250	0255	.0260	.0264	.0269	.0274	.0278	.0283
.007	.0288	.0292	.0297	.0302	.0307	.0312	.0317	.0322	.0327	.0332
.008	-0337	.0343	.0348	.0354	.0359	.0365	.0370	.0375	.0380	.0386
.009	.0391	-0397	.0402	.0407	.0412	-0417	.0422	.0427	.0432	-0437
.010	.0442	.0447	.0451	.0456	.0460	.0465	.0476	.0475	.0480	.0485
.011	.0490	.0496	.0501	.0506	.0512	.0517	.0522	-0527	.0533	.0538
.012	.0543	.0549	-0554	.0559	.0564	.0569	.0574	-0579	.0584	.0589
.013	.0594	-0600	.0605	.0611	.0616	.0622	.0627	.0633	.0638	.0643
.014	.0648	.0654	.0659	.0665	-0471	.0676	.0682	.0687	.0693	.0699
.015	.0705	.0711	.0717	.0723	-0729	.0735	.0741	.0747	.0753	.0759

SUGAR USED UP IN FERMENTATION.

Example.—Suppose the "extract gravity" is 1.0389 and the specific gravity of the alcoholic distillate is 0.9902, both at 15.6. Then 1 - 0.9902 = 0.0098, the "degree of spirit indication." From the above table the corresponding "degree of gravity lost" is found to be 0.0432.

0.0432+1.0380=1.0821, the original gravity of the wort.

The calculation in the above simplified form is accurate for normal beer wherein the free acid present, expressed as acetic, does not exceed 0.1%. In case of beer that has developed free acid much in excess of the above limit, a correction should be added to the degrees of spirit indication. This correction, which in practice it is rarely necessary to apply except in extreme cases of old or sour beer, is calculated as follows:

If a represents the grams of free acid (as acetic) in 100 cc., then the correction to be added to the spirit indication = 0.0013a - 0.00014.

Example.—Supposing the "extract gravity" to be 1.0413, the specific gravity of the alcoholic distillate to be 0.9890, and the free acid as acetic to be 0.35%. Then 1-0.989=0.0110, the degree of spirit indication.

 $0.35 \times 0.0013 - 0.00014 = 0.0003$, correction to be added to the spirit indication.

o.o110+0.0003=0.0113, corrected spirit indication.

From the above table the corresponding degrees of gravity lost are 0.0506:

0.0506 + 1.0413 = 1.0019, the original gravity of the wort.

Determination of Degree of Fermentation.—This is calculated by the formula $D = \frac{200A}{B}$, in which D = degree of fermentation, A = per cent of alcohol by weight, and B = the original extract.

Determination of Reducing Sugars.—Dealcoholize 25 cc. of the beer and make up to 100 cc. Determine reducing sugars by the Defren-O'Sullivan or Munson-Walker method, and calculate as maltose.

Determination of Dextrin.—Dilute 50 cc. of the beer to 200 cc., hydrolize by heating in a boiling water-bath for $2\frac{1}{2}$ hours with 20 cc. of hydrochloric acid (specific gravity 1.125), nearly neutralize the free acid with sodium hydroxide, make up to 300 cc., filter, and determine the dextrose by copper reduction. Multiply the amount of reducing sugars as maltose by 0.95, subtract this from the dextrose, and multiply the difference by 0.9, thus obtaining the dextrin in the beer

Determination of Glycerin.—Proceed as directed on page 703 under wine. The milk of lime is added during evaporation after the carbon dioxide has been expelled. It is advisable that the filtrate, after being evaporated to a syrupy consistency, be treated again with 5 cc. of absolute alcohol and two portions of 7.5 cc. each of absolute ether. If clear, continue as directed. If not clear, it is necessary to repeat the treatment with lime.

Determination of Total, Fixed, and Volatile Acids. — A measured volume of the beer, say 10 cc., is freed from carbon dioxide by bringing to boiling. It is then cooled and titrated with tenth-normal sodium hydroxide, using neutral litmus solution as an indicator. Each cubic centimeter of tenth-normal alkali is equivalent to 0.009 gram of lactic acid, in which the total acidity is usually expressed.

Fixed acid, also expressed as lactic, though small quantities of succinic, tannic, and malic acids are usually also present, is determined as follows: Dealcoholize a measured amount of the beer, say 10 cc., by evaporation to one-fourth its volume, dilute with water to the original volume, and titrate with tenth-normal alkali, as before.

Volatile acid is expressed as acetic, and is usually calculated by difference between total and fixed acid. Each cubic centimeter of tenthnormal alkali is the equivalent of 0.006 gram acetic acid. Determination of Proteins.—Fifty cc. of the beer are first treated with 5 cc. of dilute sulphuric acid, and concentrated by boiling to syrupy consistency. Then proceed by the Gunning method, p. 69. $N \times 6.25 =$ proteins.

Determination of Phosphoric Acid.—Unless the sample is very dark-colored, sufficiently close results can usually be obtained by direct titration of the beer itself with uranium acetate solution. For very accurate results the ash should be used. Prepare a solution of uranium acetate of such strength that 20 cc. will correspond to 0.1 gram P₂O₅. This solution is best standardized against pure, crystallized, uneffloresced, powdered hydrogen sodium phosphate, 10.085 grams of which are dissolved in water and made up to a liter. 50 cc. of this solution contains 0.1 gram phosphoric anhydride, if the salt is pure. If the solution is of proper strength, 50 cc. evaporated to dryness and ignited in a tared platinum dish should have an ash weighing 0.1874 gram. For preliminary trial about 35 grams of uranium acetate are dissolved in water, 25 cc. of glacial acetic acid, or its equivalent in weaker acid added, and the solution made up to a liter with water.

To standardize, 50 cc. of the standard phosphate solution prepared as above are heated to 90° or 100° C., and the uranium solution run in from a burette till the resulting precipitate of hydrogen uranium phosphate is complete. The end-point is determined by transferring a few drops of the solution to a porcelain plate, and touching with a drop of freshly prepared potassium ferrocyanide solution. When the slightest excess of uranium acetate has been added, a reddish-brown color is produced by the ferrocyanide. The uranium acetate solution is purposely made rather stronger than necessary at first, and by repeated trials is brought by dilution with water to the required strength (20 cc. equivalent to 50 cc. of the phosphate solution).

Fifty cc. of the beer are heated to 90° or 100° C. and titrated with the uranium acetate solution under the same conditions and in precisely the same manner as when standardizing that solution. Each cubic centimeter of the uranium acetate corresponds to 0.01% of P₂O₅.

For the phosphoric acid determination in the ash, 50 cc. of the beer are incinerated in the regular manner, and the ash moistened with concentrated hydrochloric acid. The acid is then evaporated off on the water-bath, after which the ash is boiled with 50 cc. of distilled water, and titrated with the standard uranium solution.

Determination of Carbon Dioxide.*—In the case of beer and other carbonated drinks put up in corked bottles, the carbon dioxide may be readily determined by piercing the cork with a metal champagne tap, which is connected by a flexible tube, first with a safety flask and then with an absorption apparatus somewhat after the style of that used in the determination of carbon dioxide in baking powder, the liberated carbon dioxide being absorbed for weighing in a concentrated solution of potassium hydroxide contained in a tared Liebig bulb. The beerbottle thus connected is immersed in a vessel of water, which is heated over a gas-flame, after all the carbon dioxide that will escape spontaneously has been allowed to do so. Before weighing the absorbed carbon dioxide, the beer-bottle is replaced by a soda-lime tube, and a current of air drawn through the tubes.

Beer and ale put up in bottles having patent metallic or rubber stoppers cannot thus be treated. In this case a measured quantity, say 200 cc., of the sample is transferred as quickly as possible to a large flask provided with an outlet-tube having a glass stopper, this being connected up with the safety-flask and absorption-tubes. In this case heat may be directly, though cautiously, applied to the flask containing the beer by means of a gas-flame, after all the carbon dioxide has gone over that will do so spontaneously. Exactly the same apparatus as that shown in Fig. 71 may be used to advantage for determination of carbon dioxide in beer, except that a larger distilling-flask should be used in the case of beer.

Detection of Bitter Principles.—Elaborate schemes have been worked out for the systematic treatment of beer and ale for bitter principles. Nearly all of these are complicated and somewhat unsatisfactory. The presence of alkaloids in malt liquors, deliberately introduced during the process of manufacture, is now so rare that the analyst need seldom look for them, except in cases of suspected poisoning, when the scheme of Dragendorf† or of Otto-Stas should be employed. While it is somewhat difficult to positively identify the various alkaloids, it is usually easy to prove their absence in clear solutions, if on treatment with either of the general alkaloidal reagents, Mayer's solution (Reagent No. 170), or iodine in potassium iodide (Reagent No. 143), no precipitate is formed.

It is comparatively easy to prove the mere presence or absence of hop substitutes. The bitter principle of hops is readily soluble in ether, when a sample of the beer evaporated to syrupy consistency is extracted

^{*} U. S. Dept. of Agric., Bur. of Chem., Bul. 65, p. 95; Bul. 107 (rev.), p. 92.

[†] Gerichtlich-Chemische Ermittelung von Giften, St. Petersburg, 1876.

therewith, while the bitters of quassia and aloes, common hop substitutes, are insoluble in ether. Though many varieties of bitters might be employed that are soluble in ether, the *absence* of a bitter taste from the ether extract is evidence of the absence of hops.

The most marked difference analytically between hops and their substitutes in malt liquors lies in the fact that the bitter principle of hops is completely precipitated therefrom by treatment of the beer with lead acetate (either basic or neutral), leaving no bitter taste in the filtrate after concentration, while if any of the hop substitutes are present, the concentrated filtrate from the lead acetate treatment will have a bitter taste. The excess of lead should be removed from the filtrate, before concentration and tasting, by treatment with hydrogen sulphide. If the residue from the ether or chloroform extraction of the concentrated filtrate from a beer after treatment with lead acetate be found to be bitter, there is positive evidence that a foreign substitute has been employed.

The following are characteristic reactions that may help to identify some of the common hop substitutes.*

Quassiin is readily soluble by chloroform from acid solution. If a residue containing quassiin be moistened with a weak alcoholic solution of ferric chloride and gently heated, a marked mahogany-brown coloration is produced.

On treatment of quassiin with bromine and sodium hydroxide or ammonia, a bright-yellow color is shown.

Chiretta is readily dissolved by ether from its aqueous solution. Its ether residue, when treated with bromine and ammonia, gives a straw color, slowly changing to a dull purple-brown. This is not true of its chloroform residue, so that it is not to be mistaken for quassia (Allen).

Gentian Bitter may be extracted by treatment of the acid liquor with chloroform. When the residue containing gentian bitter is treated with concentrated sulphuric acid in the cold, no color is produced, but on warming gently a carmine-red color is shown; if further treated with ferric chloride solution, a green-brown color is formed.

Aloes.—This substance is separated from beer by treating the dried residue from 200 cc. of the beer with warm ammonia, filtering, cooling, and treating the filtrate with hydrochloric acid. The resin of aloes is precipitated and collected on a filter. It is insoluble in cold water, ether,

^{*} Allen, Analyst, 12, 1887, p. 107.

chloroform, or petroleum ether, but is soluble in alcohol. It has a very characteristic odor, which serves to identify it. The hot-water solution gives a curdy precipitate on treatment with lead acetate.

Capsicin is extracted by treatment of the acid liquor with chloroform. It is recognizable by its sharp, pungent taste.

Detection of Arsenic.—By the Marsh Method.—Measure 100 cc. of the beer (freed from carbon dioxide by agitation) into a seven-inch porcelain evaporating-dish, add 20 cc. pure concentrated nitric acid, and 3 cc. pure concentrated sulphuric acid, and cautiously heat till vigorous chemical action sets in, accompanied by frothing and swelling of the beer. Turn the flame low or remove it altogether, and stir vigorously till the frothing ceases, after which the liquid may be boiled freely. At this stage transfer to a large casserole, and continue the boiling till nearly all the nitric acid is driven off. Then, holding the casserole by the handle, continue the heating till the mass chars and the fumes of sulphuric acid are given off, giving the casserole a rotary motion to prevent sputtering. The residue should be reduced to a dry, black, pulverulent char soon after the sulphuric acid fumes begin to come off freely. If still liquid, pieces of filter-paper should be stirred in while still heating, till the residue is dry, avoiding an excess of paper.

Cool, add 50 cc. of water, and remove the masses of char from the sides of the dish by the stirring-rod. Heat to boiling and filter. Use the filtrate for the Marsh apparatus, adding it gradually.

The arsenic mirror may be weighed in the usual manner, if of sufficient size.

Reinsch's Test.*—Two hundred cc. of the beer are acidified with 1 cc. of pure, concentrated, arsenic-free hydrochloric acid, and evaporated to half its volume. 15 cc. more of hydrochloric acid are then added, and a piece of pure burnished copper foil half an inch long and a quarter of an inch wide is immersed in the liquid and kept in it for an hour while simmering, replacing from time to time the water lost by evaporation. If after the lapse of an hour the copper still remains bright, no arsenic is present.

If the copper shows a deposit, remove, wash with water, alcohol, and ether, and dry. Then place the copper in a subliming-tube, and heat over a low flame. Tetrahedral crystals, apparent under the microscope, show the presence of arsenic. Blackening of the copper does not in itself prove arsenic.

^{*} Jour. Soc., Chem Ind., 20, p. 646.

Detection and Determination of Preservatives. —See Chapter XVIII. Sulphurous acid may be determined by direct titration, as in the case of wine

MALT EXTRACT.

True malt extract is a syrupy fluid having a specific gravity of from 1.3 to 1.6, and made up in accordance with the following directions of the 1880 Pharmacopœia: Upon 100 parts of coarsely powdered malt contained in a suitable vessel, pour 100 parts of water, and macerate for six hours. Then add 400 parts of water, heated to about 30° C. and digest for an hour at a temperature not exceeding 55° C. Strain the mixture with strong pressure. Finally, by means of a water-bath or vacuum apparatus, at a temperature not exceeding 55° C., evaporate the strained liquid rapidly to the consistence of thick honey.

Keep the product in well-closed vessels in a cool place.

Such an extract has a residue of at least 70%, consisting chiefly of maltose, and contains about 2% of diastase. It should furthermore be capable of converting its own weight of starch at 55° C. in less than ten minutes.

The following are analyses of three samples of pure malt extract:*

	Specific Gravity.	Alcohol.	Extract.	Free Acids as Acetic.	Volatile Acids.	Albumin- oids.	Maltose.	Dextrin.	Ash.	Phosphoric Acid.	Diastatic Action.
В	1.387 1.421 1.498	0	72.31 76.65 79.81	0.275	0.021	3.116	65.41	6.94	1.19	0.556	Complete in less than 5 min.

There are on the market many so-called malt extracts widely advertised for their tonic and medicinal virtues, having the taste and consistency of beer or ale. In fact they are virtually beer, differing therefrom mainly in respect to price. Such "malt extracts" have no diastase, and their value as nutrients depends almost entirely on their sugar content.

Harrington† has analyzed twenty-one of the best known of these alleged malt extracts, the maximum, minimum, and mean results of his analyses being as follows:

^{*} Penn. Dept. of Agric. An. Rep., 1898, p. 85.

[†] Boston Medical and Surgical Journal, Dec. 31, 1896.

	Specific Gravity.	Alcohol.	Total Residue.	Ash.
Maximum	1.0555	7-13 0-74 3-94	13.63 5.13 8.78	0.53 0.20

None of them contained any diastase, and several were preserved with salicylic acid.

DISTILLED LIQUORS.

These beverages differ from those hitherto considered, by reason of their high alcoholic content and low extract or residue. Indeed, when first distilled they are entirely without residue, but from long storage in casks, they absorb certain extractives from the wood, that impart more or less flavor as well as color.

When any fermented alcoholic infusion is subjected to distillation under ordinary circumstances, a distillate results which consists of a mixture with water of a large number of alcohols, chief among which is ethyl alcohol. The high boiling alcohols—amyl, butyl, propyl, etc., with their esters—exist in the distillate in small amount, constituting what is known as fusel oil. The various distilled liquors of commerce are made by just such a process of distillation, the product varying widely in flavor and character with the basis from which it was distilled.

The so-called pot-still (the old-fashioned copper still and worm) is well adapted for the production of potable spirits such as whiskey, brandy, gin, and rum, as these products should contain the congeneric substances which give the liquors their special character; it is not, however, suited for the manufacture of pure alcohol, because repeated distillation would be required for purification.

Now, however, by the use of improved apparatus, such as the Coffey still, involving the principle of fractional condensation, it is possible to obtain what is known as "silent spirit," or ethyl alcohol, free from fusel oil. With proper appurtenances for rectifying, one can now obtain 95% alcohol by two distillations.

Standards for Spirits.—The following are the standards adopted by the Joint Committee of the Association of Official Agricultural Chemists and the Association of State and National Food and Dairy Departments:

Distilled Spirit is the distillate obtained from a fermented mash of cereals, molasses, sugars, fruits, or other fermentable substance, and

contains all the volatile flavors, essential oils, and other substances derived directly from the material used, and the higher alcohols, ethers, acids, and other volatile bodies congeneric with ethyl alcohol produced during fermentation, which are carried over at the ordinary temperature of distillation, and the principal part of which are higher alcohols estimated as amylic.

Alcohol, Cologne Spirit, Neutral Spirit, Velvet Spirit, or Silent Spirit, is distilled spirit from which all, or practically all, of its constituents except ethyl alcohol and water, are separated, and contains not less than 94.9% (189.8 proof) by volume of ethyl alcohol.

Composition of Fusel Oil.—Fusel oil varies considerably in composition with the source from which it is derived. Amyl alcohol, being in all cases its chief constituent, is frequently known commercially as fusel oil. The alcohols found in fusel oil with their formulas, specific gravity, and boiling-points are as follows:

	Formula.	Specific Gravity.	Boiling-point.
Ethyl alcohol	C,H,OH C,H,OH C,H,OH C,H,IOH C,H,IOH	-794 -820 -803 -811	78.4° C. 97° C. 115° C. 130° C.

The following acids have been found in fusel oil, usually combined with the alcohols to form compound ethers:

Acetic	HC ₂ H ₃ O ₂	Caproic	HC ₆ H ₁₁ O ₂
Propionic	HC ₃ H ₅ O ₂	Œnanthylic	HC ₇ H ₁₃ O ₂
Butyric	HC ₄ H ₇ O ₂	Caprylic	$HC_8H_{15}O_2$
Valerianic	HC ₅ H ₉ O ₂	Pelargonic	HC,H,O,

Aging.—Freshly distilled liquors all contain notable quantities of fusel oil, which renders them harsh and unfit for use, but by the process of aging, they become in several years mellow and palatable. The chemical changes which take place during aging are discussed under whiskey.

WHISKEY.

Process of Manufacture.—Whiskey is the liquor resulting from the distillation of a fermented infusion of grain, the process being carried out in a pot-still, or some other form of still, constructed so that the resulting liquor contains not only the alcohol, but also the greater part

of the congeneric substances which are vaporized with the alcohol. The fermented infusion known as the "mash" is obtained by steeping in water the starch-containing material, usually barley, rye, corn (maize), or oats mixed with malt, and subjecting the mixture to the action of the diastase contained in the malt, in much the same manner as the mashing process in the brewing of beer, except that for whiskey the process of saccharous fermentation is carried further, with a view to obtaining a maximum yield of maltose and a minimum of dextrin. Yeast is afterwards added, and alcoholic fermentation allowed to proceed with proper precautions.

Genuine Scotch whiskey is made from malt which has been dried over peat, thus imparting a smoky taste to the liquor. Malt alone is seldom used in other whiskies; more often the grain most abundant in the locality where the whiskey is distilled forms the basis of the liquor. Bourbon whiskey (made originally in Bourbon County, Kentucky) is made from a mixture of grain, 50 to 60 per cent of which is corn, 10% malt, and the balance rye.

Corn alone mixed with malt is employed in some localities, and pure rye whiskey is made from rye and malt.

The fermented wort from whatever source obtained is subjected to distillation, purposely avoiding rectification or separation of the fusel oil and other congeneric substances which are valuable as flavors. The product of the first distillation is called "low wines," and is redistilled; the product of the second distillation is commonly divided into three fractions, of which the middle portion, or "clean spirit" is retained for the whiskey, while the first ("foreshots") and the last fraction ("faints") are mixed with the next portion of low wine to be redistilled. If the whiskey is too high in alcohol, it is diluted to the proper strength.

As new whiskey is crude and harsh in taste, it is subjected to "aging," or storing in casks for a number of years. The aging process softens and refines the flavor, but recent investigations have proved that this is not due, as formerly believed, to transformation of fusel oil into esters (ethers). The esters increase in amount during aging, as do also the acids—especially the volatile acids—the aldehydes, and the furfural. As a matter of fact, the percentage of fusel oil increases instead of diminishes on aging, due to the evaporation of water and, in a lesser degree, of alcohol through the wood; the actual amount, however, remains practically the same as at the start (see table, p. 737). When first distilled,

whiskey is perfectly colorless, but during the aging it extracts more or less color and some flavor from the casks in which it is stored. This color is especially pronounced in American whiskies, owing to the prevailing custom of charring the inside of the cask. Its flavor varies considerably with the nature of the grain used in its preparation.

- U. S. Rulings.—The following decision of President Roosevelt, based on an opinion of Attorney-General Bonaparte, was promulgated by Secretary Wilson, April 11, 1007:*
 - "Straight whiskey will be labeled as such.
- "A mixture of two or more straight whiskies will be labeled 'blended whiskey,' or 'whiskies.'
- "A mixture of straight whiskey and ethyl alcohol, provided that there is a sufficient amount of straight whiskey to make it genuinely a 'mixture,' will be labeled as compound of, or compounded with, pure grain distillate.
 - "Imitation whiskey will be labeled as such."

Joint Standards.—The following are the standards of the Joint Committee of the A. O. A. C. and the A. S. N. F. D. D.:

New Whiskey is the properly distilled spirit from the properly prepared and properly fermented mash of malted grain, or of grain the starch of which has been hydrolyzed by malt; it has an alcoholic strength corresponding to the excise laws of the various countries in which it is produced, and contains in 100 liters of proof spirit not less than 100 grams of the various substances other than ethyl alcohol derived from the grain from which it is made, and of those produced during fermentation, the principal part of which consists of higher alcohols estimated as amylic.

Whiskey (Potable Whiskey) is new whiskey which has been stored in wood not less than four years without any artificial heat save that which may be imparted by warming the storehouse to the usual temperature, and contains in 100 liters of proof spirit not less than 200 grams of the substances found in new whiskey, save as they are changed or eliminated by storage, and of those produced as secondary bodies during aging; and, in addition thereto, the substances extracted from the casks in which it has been stored. It contains, when prepared for consumption

^{*}While this decision seems to be in accord with the spirit of the Food and Drugs Act, it is liable to modification, since it is naturally not universally satisfactory. Indeed a permanent ruling, acceptable to all interests, as to what constitutes pure whiskey is extremely difficult to imagine.

as permitted by the regulations of the Bureau of Internal Revenue, not less than 45% by volume of ethyl alcohol, and, if no statement is made concerning its alcoholic strength, it contains not less than 50% of ethyl alcohol by volume, as prescribed by law.

Rye Whiskey is a whiskey in the manufacture of which rye, either in a malted condition or with sufficient barley or rye malt to hydrolyze the starch, is the only grain used.

Bourbon Whiskey is a whiskey made in Kentucky from a mash of Indian corn and rye, and barley malt, of which Indian corn forms more than 50%.

Corn Whiskey is whiskey made from malted Indian corn or of Indian corn the starch of which has been hydrolyzed by barley malt.

Blended Whiskey is a mixture of two or more whiskies.

Scotch Whiskey is whiskey made in Scotland solely from barley malt, in the drying of which peat has been used. It contains in 100 liters of proof spirit not less than 150 grams of the various substances prescribed for whiskey exclusive of those extracted from the cask.

Irish Whiskey is whiskey made in Ireland, and conforms in the proportions of its various ingredients to Scotch whiskey, save that it may be made of the same materials as prescribed for whiskey, and the malt used is not dried over peat.

U. S. P. Standards.—The requirements for whiskey are as follows: It should be at least two years old; in specific gravity it should lie between the limits of 0.945 and 0.924; its alcoholic content should be not less than 37% nor more than 47.5% by weight; the residue from 100 cc. should be not more than 0.5 gram, which should be neither sweet nor spicy, should dissolve in 10 cc. of cold water, and this solution should be colored only a pale green when treated with a drop of very dilute ferric chloride solution (a deeper color would indicate more than traces of tannin). In evaporating the liquor on the water-bath for the residue, the last traces volatilized should have an agreeable odor free from harshness, indicative of the absence of fusel oil. Its reaction should be slightly acid, but not more than 1.2 cc. of normal alkali should be required to neutralize 100 cc. of the liquor, using litmus as an indicator. If 50 cc. are shaken vigorously with 25 grams of kaolin, allowed to stand an hour and filtered, the color of the filtrate should not be much lighter than before treatment.

Composition.—Whiskey consists chiefly of alcohol and water, with relatively small amounts of fusel oil, acids, esters, aldehydes, and fur-

fural. Its extract, derived mainly from the casks in which it is stored, should consist only of small amounts of tannin, sugar, and coloring matter.

British Whiskies.—Scotch and Irish whiskies are aged in uncharred barrels, hence they are of a lighter color than the American product. Scotch whiskey is further characterized by its smoky taste, due to the peat over which it is dried. The following analyses by Vasey * illustrate the composition of Scotch and Irish whiskey of different ages, of neutral spirits used in compounding ("blending") and adulterating, and of the compounded liquors:

	Grams per 100 Liters.							
	Volatile Acids.	Esters.	Alde- hydes.	Furfural.	Fusel Oil.			
Pot-still Scotch whiskey, 8 years old	48.0	89.7	14.2	4.0	200.0			
Pot-still Scotch whiskey, 25 years old	64.8	125.1	66.1	5-4	180.0			
Irish whiskey, new		7-7	6.5	0.4	174.0			
Irish whiskey, 7 years old		20.9	11.2	3-4	204.0			
Neutral spirit for "blending"	8.4	23.8	4.9	0.4	trace			
"Blended" Scotch	39.1	106.8	14.3	3-5	108.5			
"Scotch," probably all neutral spirits	16.8	8.2	10.0	none	none			
	l		1	1	1			

It will be noted that the congeneric substances in whiskey increase on aging, although in the case of fusel oil this apparent increase is doubtless due merely to concentration dependent on evaporation. The sample of neutral spirits contained only small amounts of the congeneric substances, while the "blended" whiskies were deficient in most of these substances.

American Whiskies.—These have a deeper color than the British whiskies (due to the charred barrel) and a rich fruity flavor without the suggestion of smoke.

In the table on p. 736 are given analyses by Shepard † of fourteen leading brands, including both rye and bourbon, varying in age from four to eight years; also of two samples of neutral spirits used for compounding and adulterating.

A summary of the results obtained by Crampton and Tolman ‡ in the analysis of fourteen brands of rye and seventeen brands of bourbon whiskey at differing stages of aging appear in the table on page 737. The barrels were kept in U. S. bonded warehouses during aging, and samples

^{*} Potable Spirits, pp. 82, 83, and 87.

[†] The Constants of Whiskey, S. Dak. Food and Dairy Commission, March, 1906.

[‡] Jour. Am. Chem. Soc., 30, 1908, p. 98.

		Cent			(Grams	per 100	Liters	•		
	på	. S				Acids.					
	Age, Years.	Alcohol Per C by Volume.	Extract.	Ash.	Total.	Fixed.	Volatile.	Esters.	Aldehydes.	Furfural.	Pusel Oil.
Rye	6	50.1 50.0 49.8 50.2 49.9 50.4 50 49.8 50.1 49.8 50.1	189.8 181.5 160.4 162.1 148.5 132.7 138.6 153.7 180.0 129.3 212.0 124.5 177.2	7-3 7-4 7-3 5.8 6.4 9-7 10.0 5.1 8.0 7-2 7-3 6.5	92.0 68.4 66.8 67.1 62.4 49.2 74.8 58.8 74.4 60.9 93.0 58.2 66.5	9-3 10-2 10-2 7-5 7-5 8.6 9-9 9-9 7-2 13-5 7-2 9.0 6-3	59.1 56.6 56.9 54.9 41.7 66.2 48.9 64.5 53.7 79.5 51.0 57.5	60.7 55.9 74.8 55.9 39.6 61.6 69.6 70.8 49.3 94.0 64.0 76.6 54.6	17.5 10.0 12.0 15.0 8.0 10.5 14.0 12.5 9.5 22.5 10.0	3.2 2.4 2.6 2.6 1.0 1.3 0.7 2.5 0.8 5.0 0.5	84.9 102.6 160.4 130.9 152.0 107.4 192.7 137.1 117.0 141.7 119.5 95.3 193.6
Neutral spirits	.	95.6 94.4	10.3	0.9	7·5 6.3	I.2	6.3	15-4	2.5		30. 39.

were withdrawn at intervals of a year for eight years. As the minimum figures for certain constituents are abnormal, the next to the minimum figures are also given. It will be noted that during the first few years there was a marked increase in actual amounts of all the constituents determined, except fusel oil, over and above that due to concentration, but after three or four years the acids and esters do not materially change. The rye whiskies contained larger amounts of solids, acids, esters, etc., than the bourbons, but this was attributed to the fact that heated warehouses are used for rye, and unheated for bourbon whiskey. The authors state that the characteristic aroma of American whiskey, also the oily appearance and the "body" (solids), are due to the charred barrels.

Thirty-seven samples of whiskey, purchased by the glass from various Massachusetts saloons, were examined by the Massachusetts State Board of Health in 1894, with the following results:

	Per Cent Alcohol by Weight.	Per Cent Extract.
Maximum	45.96	1.68
Minimum.	30.70	0.08
Mean	36.51	0.50

SUMMARY OF ANALYSES OF AMERICAN WHISKIES OF DIFFERENT AGES

	İ			Grams p	er 100 I	iters of r	oo Proof	f Spirits.	
		Proof.	Color	Extract.	Acids.	Esters.	Alde- hydes.	Fur- fural.	Fusei Oil.
RYE WHISE	EY.								
New:	Average Maximum .	101. 2	0.0	13.3 30.0	4.4 72.0	16.3 21.8	5.4 15.0	1.0	90.4 161.8
	Minimum *	100.0	0.0	5.0	12.0	4-3	0.7	trace	∫ 61.8
One year old:	Average	102.5	8.8	119.7	48.6	37.0	7.0	1.8	111.6
	Maximum .	104.0	13.8 ∫ 7.2	171.0 93.0	60.5 31.1	6.8	15-5	3-3	194.0 ∫ 80.4
	Minimum *	101.0	\ 6.6	92.0	5.8	6.8	2.8	0.4	66.4
Two years old:	Average Maximum .	104.9 109.0	11.6 16.7	144.7	51.9 75.6	54.0 75.1	10.5	2.2 5-7	112.4
	Minimum *	100.0	∫ 8.8	121.0	44-3	41.5}	5-4	0.7	83.4
Three years old:		107.7	\ 8.6 13.2	94.0 171.4	11.0 62.7	31.2 J 61.5	12.5	1.5	112.7
	Maximum .	112.0	18.3	224.0	81.8	79.8	20.8	6.1	202.0
	Minimum *	104.0	11.4	145.0	52.3 16.4	47.6 34.3	6.5	0.7	79.0 60.0
Four years old:		111.8	14.0	185.0	65.9	69.3	13.9	2.8	125.1
	Maximum .	118.0	18.9 ∫11.6	238.0 156.0	83.8 58.6	89.1 57.7 \	22.1	6.7	203.5 83.8
Pinks man ald.	Minimum *	105.0	11.3	153.0	17.3 82.9	36.3 } 89.1	6.4	0.7	67.8
Eight years old:	Maximum .	1 23. 8	18.6 24.2	<i>256.0</i> 339.0	112.0	126.6	16.0 26.5	9.4	154.2 280.3
	Minimum *	112.0	{ 13.8 13.7	214.0 200.0	73·7 31·7	68.4 40.9	7-9	0.8	{ 109.0 107.1
BOURBON V	Vuicerv					İ			
New:	Average	101.0	0.0	26.5	10.0	18.4	3.2	0.7	100.9
	Maximum .	104.0	0.0	161.0	29.1	53-2	7-9	2.0	171.3
	Minimum *	100.0	0.0	4.0	12.0	13.0	1.0	trace	
One year old:	Average Maximum .	101.8 103.0	7.1	99.6	41.1 55·3	28.6 55.9	5.8 8.6	1.6 7.9	110.1
	Minimum *	100.0	∫ 5-4	61.0	24.7	17.2}	2.7	trace	∫ 58.0
Two years old:	Average	102.2	8.6	54.0 1 26. 8	7.2 45.6	40.0	8.4	1.6	108.9
,	Maximum .	104.0	11.8	214.0	61.7	59.8	12.0	9.1	197.1
	Minimum *	100.0	6.9 5.7	81.0	25.5	11.2	5.9	0.4	86.2 42.8
Three years old:		103.0	10.0	149.3	54.3	48.1	10.5	1.7	112.4
	Maximum .	106.0	13.8 ∫ 8.9		64.8 38.4	73.0	22.1	9.5	88.0
Four was model.	Minimum.*	100.0	1 7.0	90.0	32.1	12.1 } 53.5	5.9 11.0	1.9	43-5
Four years old:	Average Maximum .	104.3	10.8		58.4 73.0	80.6	22.0	9.6	123.9 237.1
	Minimum *	100.0	8.6	1	40.4	13.8	6.9	0.8	{ 95.0
Eight years old:		111.1	14.2		76.4	65.6	12.9	2.1	143.5
	Maximum .	124.0	20.9 ∫ 12.3		91.4 64.1	93.6	28.8	10.0	241.8
	Minimum *	102.0	12.3		53-7	37.7	8.7	1.0	47.6

^{*} Minimum and next to the minimum.

Seven of these samples had an excess of tannic acid, three had no tannic acid at all, and two or three had insoluble residues.

Adulteration of Whiskey.—Imitation whiskey is often concocted by diluting alcohol or neutral spirit to the proper strength, coloring with caramel, sometimes adding for body prune juice, and adding for flavor certain essential oils, such as oil of wintergreen, and artificial fruit essences, such as cenanthic and pelargonic ethers. As a rule, a small amount of pure whiskey is mixed with the artificial to give it flavor.

What has long been known as "blended whiskey" is either an imitation pure and simple, or a compound of whiskey and neutral spirits, artificially colored and flavored. According to the U. S. decisions, the term "blended whiskey" is restricted to a mixture of different kinds of genuine whiskey, colored and flavored.

Among Fleischman's recipes for "blended" whiskey is the following, which he claims to be the very lowest grade:

Spirits	_	•
Caramel	4	ounces
Beading oil	1	ounce

"Beading oil" is prepared by mixing 48 ounces oil of sweet almonds with 12 ounces C. P. sulphuric acid, neutralizing with ammonia, adding double the volume of proof spirits, and distilling. This preparation is so called because it is largely used for putting an artificial bead on cheap liquors.

A little creosote is sometimes added to give a burnt taste in semblance of Scotch whiskey. Pungent materials such as cayenne pepper are said to be used as adulterants, but no record is known of any substance being used more injurious than the alcohols. Sugar is a frequent adulterant.

Some doubt exists as to the injurious effects of fusel oil on the system. The following analyses by Ladd * show the composition of neutral spirits, and imitation whiskey consisting of neutral spirits diluted with water, colored with caramel and flavored:

^{*} N. Dak. Agric. Exp. Sta. Rep., 1906, Part II, p. 145.

	t by						Liters	•		
	er Cent				Acids.					
	Aleohol Per Volume.	Extract.	Ash.	Total.	Fixed.	Volatile.	Esters	Aldehydes	Furfural.	Pusel Oil.
Neutral spirits		2.4 366.4† 854.0†		7.2 43.2 20.4	9.0	7.2 34.2 17.4	3-5	6.0 trace trace		28.0 37.0
" " rye		456.ot	5-5	9.6		6.6	5.2			42.3

[†] Includes caramel color.

BRANDY AND COGNAC.

Brandy is the product of the distillation of fermented grape juice or wine. In a broader sense the term brandy is sometimes applied to liquor distilled from the juices of other fruits, such as apples, peaches, cherries, etc. The finest grades of brandy, such as pure cognac and armagnac (named from towns in France in which they were originally distilled). are made from choice white wine by the use of pot stills, and naturally command a high price. Brandy of a lower grade is distilled from the cheaper wines, and sometimes from the fermented marc, or refuse, of the grape, as well as from the lees and "scrapings" of the casks. The best brandies are sometimes rectified by a second distillation. Like whiskey, the fresh brandy is colorless, and would so remain if stored in glass or stone. The color is due to the wooden casks in which it is stored. Brandy consists of nearly pure alcohol and water, having a characteristic flavor. differing somewhat according to the nature and quality of the wine from which it was prepared. The chief flavor of pure cognac is due to cenanthic ether.

Composition.—Vasey gives the following analyses of cognac and of brandy adulterated with neutral spirits:

	Cognac Ten Years Old.	Brandy M	ixed wit	h Neutral Spirits.
Volatile acids	74-5			r100 liters.
Esters	109.3	32.4	"	4.4
Aldehydes	16.6	7-4	"	"
Furfural	1.7	0.6	"	"
Fusel oil	124.2	49.0	"	**

^{*} Analysis of Potable Spirits, p. 20.

Thirty-seven samples of brandy, collected from Massachusetts barrooms in 1894 and examined by the State Board of Health, showed the following results:

	Per Cent Alcohol by Weight.	Per Cent Extract.
Maximum	50.70	3.00
Minimum.	21.30	0.10
Mean	40.54	0.93

Three of these samples were artificially prepared mixtures of alcohol and water, one showed the presence of cloves, five contained tannin in excess, nine were excessively acid, and two had insoluble residues.

Joint Standards.—The following are the standards of the A. O. A. C. and the A. S. N. F. D. D.:

New Brandy is a properly distilled spirit made from wine, and contains in 100 liters of proof spirit not less than 100 grams of the volatile flavors, oils, and other substances, derived from the material from which it is made, and of the substances congeneric with ethyl alcohol produced during fermentation and carried over at the ordinary temperatures of distillation, the principal part of which consists of the higher alcohols estimated as amylic.

Brandy (Potable Brandy) is new brandy stored in wood for not less than four years without any artificial heat save that which may be imparted by warming the storehouse to the usual temperature, and contains in 100 liters of proof spirit not less than 150 grams of the substances found in new brandy, save as they are changed or eliminated by storage, and of those produced as secondary bodies during aging; and, in addition thereto, the substances extracted from the casks in which it has been stored. It contains, when prepared for consumption, as permitted by the regulations of the Bureau of Internal Revenue, not less than 45% by volume of ethyl alcohol, and, if no statement is made concerning its alcoholic strength, it contains not less than 50% by volume of ethyl alcohol as prescribed by law.

Cognac, Cognac Brandy, is brandy produced in the departments of the Charente and Charente Inferieure, France, from wine produced in those departments.

U. S. Pharmacopæia Standards.—According to the U. S. Pharmacopæia, brandy should be at least four years old; its specific gravity should be

not more than 0.941 nor less than 0.925; its alcoholic content should be from 39 to 47 per cent by weight; the residue from 100 cc. should not exceed 0.5 gram, and should dissolve readily in 10 cc. of cold water, and this solution should not be colored deeper than a pale green by the addition of dilute ferric chloride solution (absence of more than traces of tannin); the residue should not be sweet nor spicy in taste; a marked disagreeable pungent odor of fusel oil should not manifest itself on the volatilization of the last traces of alcohol in evaporating for the residue; in acidity, not more than 1 cc. of tenth-normal alkali should be required to neutralize 100 cc. of the brandy, using litmus as an indicator.

Adulteration of Brandy.—Much of the brandy sold on the market is a compound or imitation, having for its basis alcohol reduced to the requisite strength, flavored either by the admixture of real brandy, or by various preparations such, for example, as syrup of raisins, prune juice, rum, acetic ether, cenanthic ether, infusion of green walnut-hulls, infusion of bitter almond shells, catechu, balsam of Tolu, etc.

Fleischmann gives the following recipe for artificial brandy of the cheapest grade:

Spirits	45 gallons
Coloring (caramel)	6 ounces
Cognac oil	1 ounce

"Cognac oil" is made up of melted cocoanut oil 16 ounces, sulphuric acid 8 ounces, alcohol 16 ounces, mixed and distilled.

While commercial brandy often fails to meet the pharmacopœial requirements, and may contain any of the above flavoring materials, not one sample has been found among the many examined by the Massachusetts Board of Health during upwards of twenty years containing a more injurious ingredient than alcohol.

Genuine new brandy may be "aged" or "improved" for immediate use, according to Duplais, by adding to 100 liters the following:

The addition of sugar and caramel to brandy is very common. The

^{*} Brandy distilled from cherry wine.

lack of flavor resulting from the employment of "silent spirit," or from watering the product, may be compensated for by the employment of so-called cognac essences sold for the purpose, containing mixtures of the aromatic compounds named above.

RIIM.

Rum is the liquor distilled from fermented molasses or cane juice, or from the scum and other waste juices from the manufacture of raw sugar. The molasses wort is mixed with the residue from a previous fermentation and allowed to ferment for a number of days, after which it is distilled twice and stored in wood for a long time, to remove the disagreeable odor, which in the new product is especially marked. The characteristic flavor of old rum is due to a mixture of butyric and acetic ether, principally the former. Pineapples and guavas are often put in the still to impart a fruity flavor. The best varieties of rum come from Jamaica and Vera Cruz.

Composition.—The following analysis of rum is by Vasey: *

Volatile acids	28.0 gr	ams per 10	∞ liters
Esters	399.0	"	"
Aldehydes	8.4	"	"
Furfural	2.8	"	"
Fusel oil	00.6	"	"

Thirty-nine samples of rum, sold at retail in Massachusetts in 1894, were examined by the State Board of Health with the following results:

	Per Cent Alcohol by Weight.	Per Cent Extract.
Maximum. Minimum. Mean.	42.9 24.7 37.1	3-93 0-04 0-51

Of these, two samples were new rum, and several were entirely artificial.

Joint Standards.—The following are the joint standards of the A. O. A. C. and the A. S. N. F. D. D.:

^{*} Analysis of Potable Spirits, p. 85.

New Rum is properly distilled spirit made from the properly fermented clean, sound juice of the sugar cane, the clean, sound massacuite made therefrom, clean, sound molasses from the massecuite, or any sound clean intermediate product save sugar, and contains in 100 liters of proof spirit not less than 100 grams of the volatile flavors, oils, and other substances derived from the materials of which it is made, and of the substances congeneric with the ethyl alcohol produced during fermentation, which are carried over at the ordinary temperatures of distillation, the principal part of which is higher alcohols estimated as amylic.

Rum (Potable Rum) is new rum stored not less than four years in wood without any artificial heat save that which may be imparted by warming the storehouse to the usual temperature, and contains in 100 liters of proof spirit not less than 175 grams of the substances found in new rum, save as they are changed or eliminated by storage, and of those produced as secondary bodies, during aging; and, in addition thereto, the substances extracted from the casks. It contains, when prepared for consumption as permitted by the regulations of the Bureau of Internal Revenue, not less than 45% by volume of ethyl alcohol, and if no statement is made concerning its alcoholic strength, it contains not less than 50% by volume of ethyl alcohol as prescribed by law.

More or less factitious rum is sold on the market, made up of alcohol diluted to the right strength, colored with caramel, and flavored by the addition of "rum essence." Prune juice is sometimes added.

Fleischman gives the following recipe for low-grade artificial rum:

Spirits	40	gallons
New England rum	5	"
Prune juice	•	
Caramel	12	ounces
Rum essence	8	"

The "rum essence" is made up by distilling 32 ounces of a mixture of 2 ounces black oxide of manganese, 4 ounces pyroligneous acid, 32 ounces alcohol, and 4 ounces sulphuric acid. To this is added 32 ounces of acetic ether, 8 ounces of butyric ether, 16 ounces saffron extract, and $\frac{1}{8}$ ounce oil of birch.

GIN.

Gin is an alcoholic liquor, flavored with the volatile oil of juniper and sometimes with other aromatic substances, such as coriander, grains of paradise, anise, cardamom, orange-peel, and fennel. The choicest variety is known as Schiedam schnapps, named from the town of Schiedam in Holland, where there are upwards of 200 distilleries devoted to the manufacture of gin. The mash used for this variety is fermented by yeast from malted barley and rye, after which it is distilled and redistilled in pot stills with juniper berries and sometimes hops.

Juniper berries, to which the most characteristic flavor of gin is due, are dark blue in color, and possess a pungent taste. They grow on the slender evergreen shrub *Juniperus communis*. Gin differs from the other distilled liquors by being water-white. To this end it is kept in glass and not in wood.

Much of the gin of commerce is made by redistilling corn or grain whiskey with oil of juniper, and frequently one or several of the abovenamed flavoring materials. Sugar is often added, and sometimes in the cheaper productions oil of turpentine is substituted for juniper oil.

Composition.—The following analysis of unsweetened gin is by Vasey:*

Volatile acids	0.0	grams per	100 liters
Esters	37-3	"	"
Aldehydes	1.8	"	"
Furfural	0.0	"	"
Fusel oil	44.6	"	"

Thirty-three samples of gin, purchased in Massachusetts saloons and analyzed by the State Board of Health in 1894, gave the following results in per cent of alcohol by weight: Maximum 42.5, minimum 29.5, mean 38.2.

^{*} Analysis of Potable Spirits, p. 85.

METHODS OF ANALYSIS OF DISTILLED LIQUORS.

Specific gravity and alcohol are determined as described on pp. 657-677. The following methods with the exception of the qualitative test for fusel oil, Mitchell's method, and McGill's opalescence test are those of the A. O. A. C.*

Determination of Extract.—Weigh or measure (at 15.6° C.) 100 cc. of the sample, evaporate nearly to dryness on the water-bath, then transfer to a water-oven, and dry at the temperature of boiling water for 2½ hours.

Determination of Acids.—Titrate 100 cc. (or 50 cc. diluted to 100 cc. if the sample is dark in color) with tenth-normal alkali, using phenolphthalein as indicator. I cc. of tenth-normal alkali is equal to 0.006 of acetic acid.

Determination of Esters.—Dilute 200 cc. of the sample with 25 cc. of water and distil slowly into a graduated 200-cc. flask until nearly filled to the mark. Complete the volume, shake, and use aliquot portions for the determination of esters, aldehydes, and furfural.

Exactly neutralize 50 cc. of the distillate with tenth-normal alkali, using phenolphthalein as indicator, and add from 25 to 50 cc. of the tenth-normal alkali in excess of that required for neutralization. Either boil for one hour with a reflux condenser, or allow to stand overnight in a stoppered flask, and heat with a tube condenser for one-half hour below the boiling-point. Cool, and titrate with tenth-normal acid, using phenolphthalein as indicator. Multiply the number of cc. of tenth-normal alkali used in the saponification by 0.0088, thus obtaining the grams of esters calculated as ethyl acetate.

Determination of Aldehydes.—1. Reagents.—(a) Alcohol Free from Aldehydes.—Prepare by first redistilling the ordinary 95% alcohol over caustic soda or potash, then add from 2 to 3 grams per liter of m-phenylenediamine hydrochloride, digest at ordinary temperature for several days (or reflux on the steam-bath for several hours), and then distil slowly, rejecting the first 100 cc. and the last 200 cc.

(b) Sulphite-fuchsin Solution.—Dissolve 0.50 gram of pure fuchsin in 500 cc. of water, then add 5 grams of SO₂ dissolved in water, make up to a liter, and allow to stand until colorless. Prepare this solution in small quantities, as it retains its strength for only a very few days.

^{*}U. S. Dept. of Agric., Bur. of Chem., Bul. 107 (rev.), pp. 95 to 101; Circular 43.

(c) Standard Acetic Aldehyde Solution.—Prepare according to the directions of Vasey * as follows: Grind aldehyde ammonia in a mortar with ether, and decant the ether. Repeat this operation several times, then dry the purified salt in a current of air and finally in a vacuum over sulphuric acid. Dissolve 1.386 grams of this purified ammonium aldehyde in 50 cc. of 95% alcohol, to this add 22.7 cc. of normal alcoholic sulphuric acid, then make up to 100 cc. and add 0.8 cc. to compensate for the volume of the ammonium sulphate precipitate. Allow this to stand over night and filter. This solution contains 1 gram of acetic aldehyde in 100 cc. and will retain its strength.

The standard found most convenient for use is 2 cc. of this strong aldehyde solution diluted to 100 cc. with 50% alcohol by volume. One cc. of this solution is equal to 0.0002 gram of acetic aldehyde. This solution should be made up fresh every day or so, as it loses its strength.

2. Process.—Determine the aldehyde in the distillate prepared for esters. Dilute from 5 to 10 cc. of the distillate to 50 cc. with aldehyde-free alcohol (50% by volume), add 25 cc. of the fuchsin solution, and allow to stand for fifteen minutes at 15° C. The solutions and the reagents should be at 15° C. before they are mixed. Prepare standards of known strength in the same way.

Determination of Furfural.—Standard Furfural Solution.—Dissolve 1 gram of redistilled furfural in 100 cc. of 95% alcohol. This strong solution will keep. Standards are made by diluting 1 cc. of this solution to 100 cc. with 50% by volume alcohol. One cc. of this solution contains 0.0001 gram furfural.

Process.—Dilute from 10 to 20 cc. of the distillate, prepared as described under esters, to 50 cc. with furfural-free alcohol (50% by volume). To this add 2 cc. of colorless anilin and 0.5 cc. of hydrochloric acid (specific gravity 1.125), and keep for fifteen minutes in a water-bath at about 15° C. Prepare standards of known strength in the same way.

Detection of Fusel Oil.—In the process of dealcoholizing a liquor by evaporation in an open dish over the water-bath, one may readily detect fusel oil, if present, by its harsh and nauseating odor, if the nose is applied just at the moment when the last traces of alcohol are going off. At this stage any considerable trace of fusel oil will be especially apparent by the effect on the throat of the one who smells it, causing

^{*} Analysis of Potable Spirits, p. 30.

an uncontrollable desire to cough. Other ways of applying the odor test consist in pouring a small portion of the spirit into the hand, and allowing it to evaporate slowly therefrom, or in rinsing out a warm glass with the liquor, observing the odor in each case.

Goebel suggests the following test, based on the detection of the volatile acids: Agitate about 30 cc. of the liquor with 2 or 3 cc. of a dilute solution of potassium hydroxide; evaporate over the waterbath to the volume of 2 or 3 cc., cool, and to the residue add 5 or 6 cc. of concentrated sulphuric acid. If fusel oil be present, the characteristic odors of valerianic and butyric acids will be apparent.

Determination of Fusel Oil.—Allen-Marquardt Method.—Add to 100 cc. of whiskey 20 cc. of half-normal sodium hydroxide, and saponify the mixture by boiling for one hour under a reflux condenser.* Connect the flasks with a distilling apparatus, distil 90 cc., add 25 cc. of water, and continue the distillation until an additional 25 cc. is collected.

Approximately saturate the distillate with finely ground sodium chloride, and add a saturated solution of sodium chloride until the specific gravity is 1.10.

Extract this salt solution four times with carbon tetrachloride,† using 40, 30, 20, and 10 cc. respectively, and wash the carbon tetrachloride three times with 50-cc. portions of a saturated solution of sodium chloride, and twice with saturated solution of sodium sulphate. Then transfer the carbon tetrachloride to a flask containing 5 cc. of concentrated sulphuric acid, 45 cc. of water, and 5 grams of potassium bichromate, and boil for eight hours under a reflux condenser.

Add 30 cc. of water, and distil until only about 20 cc. remain; add 80 cc. of water, and distil until but 5 cc. are left. Neutralize the distillate to methyl orange, and titrate with sodium hydroxide, using phenol-phthalein as indicator. One cc. of tenth-normal sodium hydroxide is equivalent to 0.0088 gram of amyl alcohol.

Rubber stoppers can be used in the saponification and first distillation, but corks covered with tinfoil must be used in the oxidation and second distillation. Corks and tinfoil must be renewed frequently.

^{*}Or 100 cc. of the liquor may be mixed with 20 cc. of half-normal sodium hydroxide, allowed to stand overnight at room temperature, and distilled directly.

[†] Purify 5 liters of carbon tetrachloride by boiling for several hours under a reflux condenser with 200 cc. of sulphuric acid and 25 grams of potassium bichromate in 200 cc. of water; separate from the oxidizing mixture by distillation, and redistil over barium carbonate.

Tolman and Hillyer's Modification of the Allen-Marquardt Method,-Proceed with the Allen-Marquardt method to the point where the carbon tetrachloride solution of the higher alcohols is ready to be oxidized. Add 50 cc. of a solution of 200 grams of pulverized potassium bichromate in 1800 cc. of water and 200 cc. of concentrated sulphuric acid, very carefully measured with pipette or burette, and start the eight-hour oxidation. Take great care to prevent any isolation of spots of bichromate on the flask during the oxidation. Decomposition of the bichromate from overheating can best be prevented by slow boiling over several thicknesses of asbestos board. After the oxidation is complete, separate the bichromate solution from the carbon tetrachloride in a separatory funnel, care being taken to wash the carbon tetrachloride free from bichromate. Make up the bichromate solution to 500 cc. Place 200 cc. of this solution in a liter flask, add 20 cc. of concentrated hydrochloric acid, 100 cc. of potassium iodide solution (1:1), and 50 cc. of approximately three-fourths normal thiosulphate not standardized. Make this last addition by means of a burette. (If a high content of fusel oil is present, 50 cc. of thiosulphate may be excessive and a smaller amount should be used, the same quantity being added to the sample and to the blank.) Run blanks containing exactly the same amount of reagents with each series, and treat them in the same way, starting them at the point where the carbon tetrachloride is washed with sodium chloride. The titration of this blank, to which has been added exactly the same amount of three-fourths normal thiosulphate, gives the value of the bichromate solution. The difference in cubic centimeters of tenthnormal thiosulphate used in titrating the blank and the samples gives the amount of bichromate reduced by the higher alcohols. This difference in cubic centimeters of tenth-normal thiosulphate multiplied by the factor 0.001773 gives grams of higher alcohols present.

Mitchell's Method.*—This method is more rapid than the Allen-Marquardt method and gives more nearly the true amount of fusel oil. Saponify, distil, shake with sodium chloride, and extract with carbon tetrachloride, as in the Allen-Marquardt method. To the carbon tetrachloride extract, contained in the separatory funnel, add 10 cc. of potassium hydroxide solution (1:1). Cool the mixture in ice-water to approximately 0° C. Similarly cool 100 cc. of a solution of potassium permanganate solution (20 grams to the liter), accurately measured in

^{*}A. O. A. C. Proc., 1908, U. S. Dept. of Agric., Bur. of Chem., Bul. 122, p. 199.

a flask. To the contents of the separatory funnel add the bulk of the permanganate solution, but without rinsing, retaining the residue to be added at a later stage. Remove the mixture from the bath, and shake vigorously for five minutes; set aside for thirty minutes, with occasional shaking, permitting the liquid to warm to room temperature (20 to 25° C.)

Accurately measure into a liter Erlenmeyer flask 100 cc. of a solution of hydrogen peroxide slightly (about 2%) stronger than the permanganate solution, acidulate with 100 cc. of an approximately 25% sulphuric acid solution, and slowly add the contents of the separatory funnel with constant shaking, keeping the acid solution constantly in excess. Rinse the separatory funnel and the flask containing the residue of permanganate with water and add to the peroxide solution. Finally titrate the excess of hydrogen peroxide with standard potassium permanganate solution (10 grams to the liter).

Run a blank determination, using the same amounts of the stronger permanganate, potassium hydroxide, hydrogen peroxide, and sulphuric acid solutions, and titrating the residual peroxide with the standard potassium permanganate as before.

The difference in the amounts of permanganate consumed, in grams, times 0.696, gives the amount of amyl alcohol.

Detection of Methyl Alcohol.—Leach and Lythgoe Immersion Refractometer Method.*—Determine at 20° C. the refraction of the distillate obtained in the determination of alcohol by the immersion refractometer. If on reference to the table the refraction shows the percentage of alcohol agreeing with that obtained from the specific gravity, it may be safely assumed that no methyl alcohol is present. If, however, there is an appreciable amount of methyl alcohol, the low refractometer reading will at once indicate the fact. If the absence from the solution of other refractive substances than water and the alcohols is assured, this qualitative test by difference in refraction is conclusive.

The addition of methyl to ethyl alcohol decreases the refraction in direct proportion to the amount present; hence the quantitative calculation is readily made by interpolation in the table, using the figures for pure ethyl and methyl alcohol of the same alcoholic strength as the sample.

Example.—Suppose the distillate made up to the original volume of the measured portion taken for the alcohol determination has a

^{*} Jour. Am. Chem. Soc., 27, 1905, p. 964.

specific gravity of 0.9736, corresponding to 18.38% alcohol by weight, and has a refraction of 35.8 at 20° C. by the immersion refractometer; by interpolation in the refractometer table the readings of ethyl and methyl alcohol corresponding to 18.38% alcohol are 47.2 and 25.4, respectively, the difference being 21.8; 47.2-35.8=11.4; (11.4 ÷ 21.8) 100=52.3, showing that 52.3 of the alcohol present is methyl alcohol.

SCALE READINGS ON ZEISS IMMERSION REFRACTOMETER AT 20° C., CORRESPONDING TO EACH PER CENT BY WEIGHT OF METHYL AND ETHYL ALCOHOLS.

Per Cent	Sca Read		Per Cent	Sca Read		Per Cent	Read	ale lings.	Per Cent		ale lings.
Alcohol by Weight.	Methyl Al- cohol.	Ethyl Al- cohol.	Alcohol by Weight.	Methyl Al- cohol.	Ethyl Al- cohol.	Alcohol by Weight.	Methyl Al- cohol.	Ethyl Al- cohol.	Alcohol by Weight.	Methyl Al- cohol	Ethyl Al- cohol.
0	14.5	14-5	26	30.3	61.9		39-7	91.1	76	29.0	101.0
I	14.8	16.0		30.9	63.7		39.6	91.8	77	28.3	100.9
2	15.4	17.6	28	31.6	65.5		39.6	92.4	78	27.6	100.9
3	16.0	19.1	29	32.2	67.2		39-5	93.0	79	26.8	100.8
4	16.6	20.7	30	32.8	69.0	55	39-4	93.6	8o-	26.0	100.7
5 6	17.2	22.3	31	33-5	70.4	56	39.2	94.1	81	25.1	100.6
	17.8	24.1	32	34.1	71.7	57	39.0	94.7	82	24.3	100.5
7 8	18.4	25.9		34.7	73.1		38.6	95.2	83	23.6	100.4
	19.0	27.8		35-2	74-4		38.3	95.7	84	22.8	100.3
9	19.6	29.6	35	35.8	75.8	60	37-9	96.2	85	21.8	100.1
10	20.2	31.4	36	36.3	76.9		37-5	96.7	86	20.8	99.8
II	20.8	33.2	37	36.8	78.0		37.0	97.1	87.	19.7	99-5
12	21.4	35-0	38	37-3	79.1	63	36.5	97.5	88	18.6	99.2
13	22.0	36.9		37-7	80.2	64	36.0	98.0	89	17.3	98.9
14	22.6	38.7	40	38.1	81.3	65	35-5	98.3	90	16.1	98.6
15	23.2	40.5	41	38.4	82.3	66	35.0	98.7	91	14.9	98.3
ıŏ	23.9	42.5	42	38.8	83.3	67	34-5	99.i	92	13.7	97.8
17	24.5	44-5	43	39.2	84.2	68	34.0	99-4	93	12.4	97.2
18	25.2	46.5	44	39-3	85.2	69	33-5	99.7	94	11.0	96.4
19	25.8.	48.5	45	39-4	86.2	70	33.0	100.0	95	9.6	95-7
20	26.5	50.5	46	39.5	87.0	71	32.3	100.2	96	8.2	94.9
21	27.1	52-4		39.6	87.8	72	31.7	100.4	97	6.7	94.0
22	27.8	54-3		39-7	88.7		31.1	100.6	98	5.1	93.0
23	28.4	56.3		39.8	89.5	74	30.4	100.8	99	3-5	92.0
24	29. I	58.2		39.8	90.3		29.7	101.0	100	2.0	91.0
25	29.7	60.1	1	1						l	
	<u> </u>		·			}	<u> </u>	<u> </u>	<u> </u>	I	<u> </u>

Trillat Method.*—To 50 cc. add 50 cc. of water and 8 grams of lime, and fractionally distil by the aid of Glinksy bulb tubes. Dilute the

^{*} A. Trillat, Analyst, 24, 1899, pp. 13, 211-212.

first 15 cc. of the distillate to 150 cc., mix with 15 grams of potassium bichromate and 70 cc. of sulphuric acid (1:5), and allow to stand for one hour with occasional shaking.

Distil, reject the first 25 cc., and collect 100 cc. Mix 50 cc. of the distillate with 1 cc of rectified dimethyl-anilin, transfer to a stout, tightly-stoppered flask, and keep on bath at 70 to 80° C. for three hours with occasional shaking. Make distinctly alkaline with sodium hydroxide, and distil the excess of dimethyl-anilin, stopping the distillation when 25 cc. have passed over.

Acidify the residue in the flask with acetic acid, shake, and test a few cc. by adding four or five drops of water with lead dioxide in suspension (1 gram in 100 cc.). If methyl alcohol be present, a blue coloration occurs which is increased by boiling.

Note.—Ethyl alcohol thus treated yields a blue coloration, changing immediately to green, afterwards to yellow, and becoming colorless when boiled.

Riche and Bardy Methoa.*—The following method for the detection of methyl alcohol in commercial spirit of wine depends on the formation of methyl-anilin violet:

Place 10 cc. of the sample, previously rectified over potassium carbonate if necessary, in a small flask with 15 grams of iodine and 2 grams of red phosphorus. Keep in ice-water for from ten to fifteen minutes until action has ceased. Distil on a water-bath the methyl and ethyl iodides formed into about 30 cc. of water. Wash with dilute alkali to eliminate free iodine. Separate the heavy oily liquid which settles, and transfer to a flask containing 5 cc. of anilin. The flask should be placed in cold water, in case the action should be violent, or, if necessary, the reaction may be stimulated by gently warming the flask. After one hour boil the product with water, and add about 20 cc. of a 15% solution of soda; when the bases rise to the top as an oily layer, fill the flask up to the neck with water, and draw them off with a pipette. Oxidize 1 cc. of the oily liquid by adding 10 grams of a mixture of 100 parts of clean sand, 2 of common salt, and 3 of cupric nitrate; mix thoroughly, introduce into a glass tube, and heat to 90° C. for eight or ten hours. Exhaust the product with warm alcohol, filter, and make up with alcohol to 100 cc. If the sample of spirits be pure, the liquid is of a red tint, but in the presence of 1% of methyl alcohol, it has a distinct violet shade; with

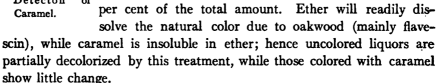
^{*} Allen's Commercial Organic Analysis, 3d ed., I, p. 80.

2.5% the shade is very distinct, and still more so with 5%. To detect more minute quantities of methyl alcohol, dilute 5 cc. of the colored liquid to 100 cc. with water, and dilute 5 cc. of this again to 400 cc. Heat the liquid thus obtained in porcelain, and immerse a fragment of white merino (free from sulphur) in it for half an hour. If the alcohol be pure, the wool will remain white, but if methylated, the fiber will become

violet, the depth of tint giving a fair approximate indication of the proportion of methyl alcohol present.

Detection of Caramel.—Crampton and Simon's Method.*—Evaporate 50 cc. of the liquor nearly but not quite to dryness in an evaporating-dish on the water-bath. Wash with water into a 50-cc. graduated glass-stoppered flask, add 25 cc. of absolute alcohol, and fill to the mark with water. Shake, and transfer 25 cc. of the solution to a separatory funnel of the type presented in Fig. 116, the stem of which terminates in a 25-cc. graduated bulb pipette, provided with a stop-cock as shown.

Add 50 cc. of ether, and shake carefully at intervals during half an hour. After complete separation, make up the lower aqueous layer with water to the 25-cc. mark, which may be done by siphoning it in through a rubber tube from an elevated flask, controlling the supply by the stop-cock. Shake the separatory funnel, and again allow the layers to separate, draw off the aqueous layer, and compare with the color of the original liquor. Express the amount of color removed as per cent of the total amount. Ether will readily dissolve the natural color due to oakwood (mainly flave-



Amthor Test, Modified by Lasché.†—Add 10 cc. of paraldehyde to 5 cc. of the sample contained in a test tube and shake. Add absolute alcohol, a few drops at a time, shaking after each addition until the mixture becomes clear. Allow to stand. Turbidity after ten minutes is an indication of caramel.



Fig. 116.—Separatory Funnel for Detecton of Caramel.

^{*} Jour. Am. Chem. Soc., 22 1900, p. 810. † The Brewer Distiller, May, 1903.

Determination of Water-insoluble Color in Whiskies.—Evaporate 50 cc. of the sample just to dryness. Take up with cold water, using approximately 15 cc., filter, and wash until the filtrate amounts to nearly 25 cc. To this filtrate add 25 cc. of absolute alcohol or 26.3 cc. of 95% by volume alcohol, and make up to 50 cc. by the addition of water. Mix thoroughly and compare in a colorimeter with the original material. Calculate the per cent of color insoluble in water from these readings.

Determination of Color Insoluble in Amyl Alcohol.—Modified Marsh Test.—Evaporate 50 cc. of the whiskey just to dryness on the steambath. Add 26.3 cc. of 95% alcohol to dissolve the residue. Transfer to a 50-cc. flask and make up to volume with water to obtain a uniform alcohol concentration. Place 25 cc. of this solution in a separatory funnel, and add 20 cc. of the Marsh reagent, shaking lightly so as not to form an emulsion. (This reagent consists of 100 cc. of pure amyl alcohol, 3 cc. of syrupy phosphoric acid, and 3 cc. of water; shake before using.) Allow the layers to separate, and repeat this shaking and standing twice again. After the layers have clearly separated, draw off the lower or watery layer which contains the caramel into a 25-cc. cylinder, and make up to volume with 50% by volume alcohol. Compare this solution in a colorimeter with the untreated 25 cc. Calculate the result of this reading to the per cent of color insoluble in amyl alcohol.

Opalescence in Diluted Alcohol Distillate.—McGill* has shown that in the case of liquors made from thoroughly rectified grain spirit, there is little or no opalescence produced when the alcoholic distillate (i.e., that used in determining the alcohol) is diluted with an equal volume of water, while in the case of liquors distilled from alcoholic infusions without rectification, the opalescence is marked. He ascribes the opalescence to the presence of minute amounts of volatile oils present in wine marc, grains, and other sources of these liquors, soluble in strong, but insoluble in dilute alcohol. Whether due to this or to the separation of minute traces of fusel oil on dilution, the presence or absence of turbidity certainly furnishes a rough distinguishing test, indicating in some cases the exclusive use of rectified spirit.

^{*} Bul. 27, Canadian Inland Rev. Dept.

LIQUEURS AND CORDIALS.

These are manufactured beverages, usually high in alcohol and sugar, flavored with a wide variety of aromatic herbs or essences, and often strongly colored. Red colors most frequently used for this purpose are cochineal, cudbear, and red sandal and Brazil woods; for yellow colors, caramel and saffron-yellow are employed; for blue, indigo; and for green, chlorophyll and malachite green.

Some of the oldest of the liqueurs, such as chartreuse and bénédictine, derive their names from certain monasteries of Europe, in which they have been made for many years.

Absinthe is one of the best-known cordials, made by redistilling 40% alcohol in which wormwood, anise, sweet flag, and marjoram leaves have been macerated. Sometimes coriander and fennel are also used. It is highly intoxicating.

Curaçoa is made by distilling dilute spirits in which Curaçoa orangepeel,* cinnamon and often other spices have been soaked, and by adding sugar to the resulting liqueur.

De Brevans gives the following recipe for curaçoa:

Rasped skins of	18	or 20	oranges
Cinnamon	4	grams	
Mace	2	66	
Alcohol (85%)	5	liters	
White sugar	•		

Macerate for fourteen days, distill without rectification, and color with caramel.

Angostura owes its flavor to Angostura bark and various spices.

Maraschino had originally for its basis the fermented juice of the sour Italian cherry, to which honey was added. It is more commonly made by distilling a mixture in alcohol of ripe wild cherries, raspberries, cherry leaves, peach nuts, and orris. Finally sugar is added.

Chartreuse and Bénédictine contain much sugar, and are flavored with the volatile oils of angelica, hyssop, nutmeg, and peppermint.

Noyau, or Crême de Noyau, is a preparation distilled from brandy, bitter almonds, mace and nutmeg. Sugar and coloring matter, usually pink, are added to the final product.

^{*} This is a very rare and highly prized orange, growing in the island of Curaçoa.

Crême de Menthe, according to De Brevans, is made by distilling a mixture of

Peppermint	600	grams
Balm	40	"
Sage	IO	"
Cinnamon		66
Orris root		"
Ginger	15	"
Alcohol (80%)	-	cc.

producing finally 10 liters of the liquor, after 3750 grams of white sugar have been introduced.

The better grades of crême de menthe were formerly colored with an alcoholic solution of chlorophyll, derived by macerating bruised green leaves of various plants with alcohol, but at present, coal-tar dyes are used. Frequently the desired shade is secured by mixing a green (e.g., Light Green S.F.), a blue-green (e.g., Malachite Green), or a blue (e.g., Indigo Carmine) with a yellow color.

The following analyses, due to König, show the chemical composition of the best-known cordials:

	Specific Gravity.	Alcohol by Vol- ume.	Alcohol by Weight.	Extract.	Cane Sugar.	Other Extrac- tives.	Ash.
Absinthe	0.9116	58.93		0.18		0.32	
Bénédictine	1.0709	52	38.5	36.∞	32-57	3.43	0.043
Ginger	1.0481	47-5	36.0	27.79	25.92	1.87	0.141
Crême de menthe	1.0447	48.0	36.5	28.28	27.63	0.65	0.068
Anisette de Bordeaux	1.0847	42.0	30.7	34.82	37-44	0.38	0.040
Curaçoa	1.0300	55.0	42.5	28.60	28.50	0.10	0.040
Kümmel	1.0830	33-9	24.8	32.02	31.18	0.84	0.058
Angostura	0.9540	49-7		5.85	4.16	1.60	
Chartreuse	1.0799	43.18		36.11	34-35	1.76	

Analysis of Cordials and Liqueurs.—The character of the essences and flavoring principles used in these beverages is so widely varied that no regular systematic plan for identifying them can be made applicable to all cases. The senses of smell and taste are most useful, both when applied directly to the liqueur itself and to the dry extract, for suggestions as to the main ingredients employed. Coloring-matters, sugars, acids, and alcohol are determined as with other liquors, except that in the case of alcohol all volatile oils must first be separated out by treatment with magnesia, as directed for alcohol in lemon extract. Presence of volatile

oils is shown, if on treatment of a few cubic centimeters of the sample in a test-tube with water a precipitate is formed.

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(See also References on Leavening Materials, page 275.)

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CHAPTER XVI.

VINEGAR.

VINEGAR is the product formed by the acetic fermentation of an alcoholic liquid under the influence of the organism mycoderma acett, existing in the "mother-of-vinegar." While vinegar may be made directly from a dilute solution of pure alcohol, it is more often obtained from fruit juice, wine, or other saccharine liquid that has first undergone alcoholic fermentation.

Of the following equations, (1) and (2) illustrate the processes of inversion and alcoholic fermentation respectively, while (3) and (4) show the double process of acetic fermentation, wherein the alcohol is oxidized, first to acetaldehyde and finally to acetic acid:

$$\begin{array}{c} C_{12}H_{22}O_{11}+H_2O=2C_6H_{12}O_6;\\ Cane sugar & Invert sugar \\ \\ C_6H_{12}O_6=2C_2H_6O+2CO_2;\\ Invert sugar, & Alcohol \\ Invert sugar, & Invert sugar \\ Invert sugar \\ Invert sugar$$

In addition to the acetic acid, its chief active principle, vinegar usually contains traces of other organic acids free or combined, small amounts of alcohol, aldehyde, sugar, glycerin, coloring matter, aromatic ethers, and mineral salts, its extract varying considerably with the source from which the vinegar was obtained.

Varieties.—The principal varieties of vinegar are the following: Cider vinegar, wine vinegar, malt or beer vinegar, spirit vinegar, glucose vinegar, molasses vinegar, and wood vinegar, the three last being more frequently used as adulterants of the others.

Manufacture of Vinegar.—Cider vinegar, the principal variety used in the United States and Canada, was formerly made almost entirely by the slow process of cask fermentation, the fresh cider being allowed to undergo both alcoholic and acetic fermentation in barrels with open bung-holes in a warm cellar, or exposed to the sun. Two or three years are required for this process. Sometimes fresh cider is added to the barrels at regular intervals of two or three weeks, thus causing a series of progressive fermentations. The acetic fermentation is hastened by adding old vinegar. or mother-of-vinegar to the cider. While farmers and some manufacturers still continue to make cider vinegar by the slow process, the quick or "generator" vinegar process is now much used for cider vinegar. though originally intended and almost exclusively used in the manufacture of malt, beer, and spirit vinegar. This process requires only two or three days for complete acetification. In the quick process, the cider or other alcoholic liquor is allowed to percolate slowly through beechwood shavings or birch twigs, held in a cask known as a generator, provided with a perforated, false bottom, the shavings or twigs being previously saturated with old vinegar, and a current of air being passed up through them.

The alcoholic liquid from which genuine malt vinegar is made is derived from the wort obtained by mashing malt, or a mixture of malt and barley. Spirit vinegar is derived from diluted whiskey, brandy, or grain alcohol. Wine vinegar is made by allowing the wine to stand over wine lees for a time, after which it is clarified by passing through beech shavings, and subjected to progressive acetification in large open oak casks, to which the wine is added, the vinegar being drawn off in much the same manner as the slow-process cider vinegar.

CHARACTERISTICS AND COMPOSITION OF THE VARIOUS VINEGARS.—Cider Vinegar is brownish yellow in color, and possesses an odor of apples. It is chiefly distinguished from other vinegar by the large amount of malic acid normally present, by the character of its sugars, and by the predominance of potash in the ash. Its specific gravity varies from 1.013 to 1.015. Its acidity varies from 3 to 6 per cent, and its solids from 1½ to 3 per cent. Cider vinegar under polarized light is always lævo-rotary.

The following are summarized data of analyses made by H. C. Lythgoe in the writer's laboratory of twenty-two samples of cider vinegar of known purity:

			Acetic Total Ash. Alkalin-	P ₂ O ₅ in A	Ash of 100 Vinegar.				
			Acid.	Solids.		Ash. I	Soluble (mgr.).	Insoluble (mgr.).	
Maximum Minimum. Average.			5.86 3.92 4.84	3.20 1.84 2.49	0.42 0.20 0.34	36.1 22.2 29.7	31.7 12.1 19.2	31.5 6.5 15.6	
	Before	After Inversion.	Polariza- tion, Degrees Ventzke 200-mm. Tube.	Malic Acid.	Per Cent Ash in Total Solids.	Per Cent Reducing Sugars in Total Solids.	Ratio of Soluble to Total P ₂ O ₈ .	Alkalin- ity of 1 Gram of Ash, cc. Nah, cc.	
Maximum Minimum Average	0.51 0.15 0.25	0.53 0.15 0.25	-3.6 -0.3 -1.3	0.16 0.08 0.11	19.0 10.0 13.8	16.6 7·3 10.7	66.9 50.0 56.3	125.0 69.0 90.0	

¹ Number of cubic centimeters of tenth-normal acid to neutralize the ash of 100 grams of vinegar.

Twenty-two samples of pure cider vinegar were analyzed by A. W. Smith * with the following results:

	Acetic Acid.	Total Solids.	Ash.	Alkalinity of Ash.	Soluble P ₂ O ₈ .	Insoluble P ₂ O ₅ .	Total P ₂ O ₅ .
Maximum	3-24	4-45 2.00 2.83	0.51 0.31 0.39	55.2 28.4 38.8	22.7 13.6 19.1	19.4 4.2 10.1	39.0 19.8 28.6

Number of cubic centimeters of tenth-normal acid required to neutralize the ash from 100 grams of vinegar.

The composition of cider vinegar ash is found by Doolittle and Hess † to be as follows:

Calcium oxide	CaO	3.4 to 8.21
Magnesium oxide	MgO	1.88 " 3.44
Potassium oxide	K ₂ O	46.33 " 65.64
Sodium oxide	Na ₂ O	None
Sulphuric anhydride	SO ₃	4.66 to 16.29
Phosphoric anhydride	P ₂ O ₅	3.29 " 6.66
Iron oxide	Fe ₂ O ₃	None " trace
	CO ₂ and loss	0.00 " 40.44

Wine Vinegar is light yellow if made from white wine, and red if from red wine. The former is the highest prized. Wine vinegar varies in specific

^{*} Jour. Am. Chem. Soc., 20 (1898), p. 6.

[†] Ibid., 22 (1900), p. 220.

gravity from 1.0129 to 1.0213, and contains from 6 to 9 per cent of acetic acid. It is characterized chiefly by the bitartrate of potassium (cream of tartar) which true wine vinegar always possesses. Free tartaric acid is also usually present. Wine vinegar is the principal vinegar of France and Germany. In the United States the term white wine vinegar is usually applied to distilled or spirit vinegar, which is much cheaper than the real wine vinegar and altogether inferior to it.

Wine vinegar is slightly lævo-rotary with polarized light.

The composition of genuine white wine vinegar is shown by the following summary of the analyses of twenty-two samples, made in the Municipal Laboratory of Paris:

•.	Specific Gravity.	Total Solids.	Sugar.	Bitartrate of Potash.	Ash.	Acidity (as Acetic).
Maximum Minimum Mean	1.0213	3.19	0.46	0.36	0.69	7.38
	1.0129	. 1.38	0.06	0.07	0.16	4.44
	1.0175	1.93	0.22	0.17	0.32	7.38

Weigmann gives the following average of analyses of red wine vinegar:

Specific Gravity.	Acetic Acid.	Total Tartaric Acid.	Free Tartaric Acid.	Cream of Tartar.	Alcohol.	Extract.	Gly- cerin.	Ash.	Phos- phoric Acid.
1.0143	7 - 79	0.216	0.006	0.057	1.19	0.863	0.141	0.118	0.012

Malt or Beer Vinegar is of a brown color, and its odor is suggestive of sour beer. It varies in specific gravity from 1.015 to 1.025; its acidity is about the same as cider vinegar, but the extract is much larger, varying from 4 to 6 per cent. Malt vinegar contains considerable nitrogenous matter, and notable quantities of phosphates, dextrin, and maltose. It contains no cream of tartar. Malt vinegar is largely used in Great Britian.

Hehner gives the following data of the analyses of seven samples of vinegar undoubtedly made from malt only.*

	Acidity.	Total Solids.	Ash.	Phosphoric Anhydride.	Alkalinity (Na ₂ CO ₃).
Maximum Minimum Mean	6.48	4.23	0.47	.13	.089
	2.88	1.68	0.22	.067	.017
	4.23	2.70	0.34	.105	.024

^{*} Analyst, 16, p. 82. See also Analyst, 18, p. 240.

Allen gives the results of the analyses of three samples of genuine vinegar brewed from a mixture of malted and unmalted barley as follows:*

	Specific Gravity.	Acetic Acid.	Total Solids.	Ash.	Alkalinity as K ₂ O.	Phos- phoric Acid.	Nitrogen.	Albumin- oids.
3		6.39 5.26 4.86	2.67 3.96 2.31	0.34 0.40 0.47	0.091 0.118	0.077 0.093 0.057	.099 .095 .099	.624 .598 .624

Distilled, Spirit, or Alcohol Vinegar.—This vinegar, being made from diluted alcohol, is nearly colorless, unless artificially colored, as it often is, with caramel. As stated on page 762, the "white wine" vinegar (incorrectly so-called) commonly sold in the United States is of this class. Its specific gravity ranges from 1.008 to 1.013. Spirit vinegar contains from 3 to 10 per cent of acetic acid. Its content of total solids is insignificant, and it contains only traces of ash. It always contains non-acetified alcohol and aldehyde. It has no optical activity with polarized light.

Twelve samples of alcohol vinegar analyzed in the Municipal Laboratory of Paris gave the following results:

	Specific Gravity.	Total Solids.	Sugar.	Ash.	Acidity.
Maximum	1.0131 1.0082 1.0100	0.16 0.07 0.35	Trace	.09 .04 Trace	7-98 4-98 6-34

Glucose Vinegar is made from the acetification of alcohol, obtained from the fermentation of commercial glucose. This vinegar usually possesses the odor and taste of fermented starch. It is low in total solids, the extract consisting almost entirely of untransformed glucose, and the vinegar therefrom contains all the ingredients of the product from which it was made, viz., dextrin, maltose, and dextrose, as well as sulphate of calcium. It is decidedly dextro-rotary with polarized light both before and after inversion.

Molasses Vinegar.—This is largely the product of the acetic fermentation of sugar-house wastes, and sometimes of the accidental acetic fermentation of molasses itself, after it has undergone alcoholic fermentation for the manufacture of rum. This variety of vinegar is sometimes

used as an adulterant of cider vinegar. With polarized light molasses vinegar is dextro-rotary before, and lævo-rotary after inversion.

Wood Vinegar is prepared by the purification of pyroligneous acid, which may be accomplished by saturating the crude acid with lime or soda, adding hydrochloric or sulphuric acid, and distilling. It is further purified by redistillation with potassium bichromate, and filtration through bone-black. Acetic acid is sometimes added to impart flavor.

The extract and ash of wood vinegar are very small. Its specific gravity averages 1.007 according to Blyth. Empyreumatic or tarry products are nearly always present in vinegar of this class.

ANALYSIS OF VINEGAR.

Specific Gravity.—This is obtained either with the hydrometer, pycnometer, or Westphal balance.

Determination of Extract or Total Solids.—Weigh 5 grams of the sample in a tared platinum dish, and evaporate to dryness over the live steam of a boiling water bath, keeping the dish thereon for two hours. Cool in a desiccator and weigh.

Determination of ash.—Transfer the dish containing the last residue or extract to a music, and burn at a low red heat to an ash, or the ignition may be accomplished with care over a direct slame turned low. Cool the dish and weigh.

Determination of Solubility and Alkalinity of the Ash.—Smith's Method.*—Twenty-five cc. of the vinegar are evaporated to dryness in a tared platinum dish, ignited, cooled, and the ash weighed. The ash is then repeatedly extracted with hot water by washing into a Gooch crucible provided with a layer of asbestos (previously ignited in the crucible, cooled, and weighed) or upon an ash-free filter. Dry the Gooch or filter, ignite, cool, and weigh the insoluble ash. The aqueous extract is titrated directly with tenth-normal acid, using methyl orange as an indicator, or treated by adding an excess of tenth-normal hydrochloric acid, boiling and titrating back with tenth-normal sodium hydroxide, using phenolphthalein. Express the alkalinity in terms of 100 grams of the vinegar, by multiplying by 4 the number of cubic centimeters of acid required to neutralize.

Determination of Phosphoric Acid.†—Extract repeatedly the insoluble ash as obtained in the preceding section with hot water acidulated with nitric acid, and acidify with nitric acid the neutralized solution of the

^{*} Jour. Am. Chem. Soc., 20, p. 5.

[†] U. S. Dept. of Agric., Bur. of Chem., Bul. 46, p. 12.

765

soluble ash. Add to each solution 15 grams of ammonium nitrate, heat to boiling, and precipitate the phosphoric acid with 50 cc. of ammonium molybdate (reagent No. 53). Digest for an hour at a temperature of about 65°, filter, and wash with cold water. Dissolve the precipitate on the filter with ammonia and hot water, and wash into a beaker to a bulk of not more than 100 cc. Nearly neutralize with hydrochloric acid, cool, and add slowly magnesia mixture (reagent No. 164) drop by drop while stirring vigorously. After fifteen minutes add 30 cc. of ammonia (specific gravity 0.96), let stand for at least two hours, filter on a Gooch crucible, wash with 2.5% ammonia till practically free from chlorides, ignite, and weigh as Mg₂P₂O₇. Express results in terms of milligrams of phosphoric anhydride in the soluble and insoluble vinegar ash from 100 grams of vinegar.

Phosphoric acid in the soluble and insoluble ash may be conveniently determined also by the uranium acetate method, page 725.

Determination of Nitrogen.—Concentrate from 50 to 100 cc. of vinegar to a syrupy consistency, and proceed as directed under the Kjeldahl or Gunning method, page 69.

Determination of Total Acidity.—Six cc. of vinegar are carefully measured from a pipette into a white porcelain dish and diluted with water. Using phenolphthalein as an indicator, titrate with tenth-normal sodium hydroxide. The number of cubic centimeters of the latter required to neutralize, divided by 10, expresses the acidity in terms of percentage of acetic acid.

Approximate Determination of Vinegar Acidity by Lime Water.—It has generally been considered difficult for vinegar dealers and others who desire to estimate the acidity of their vinegar to do this themselves, in that it has been necessary to obtain for the purpose a carefully standardized alkaline solution, the exact strength of which it is impossible for them to determine.

It has been found that very satisfactory, though of course not absolutely accurate, results may be obtained by the use of ordinary lime water, which any one may easily prepare by making a saturated solution of ordinary air-slaked lime. The strength of such a solution is very nearly constant, and has been found to be about $\frac{1}{2}$ of the normal. If, therefore, it is not easy to obtain exactly normal or tenth-normal alkali, approximate figures may be obtained by employing such a saturated lime water. If 2.75 cc. of vinegar are titrated with lime water contained in a burette, using phenolphthalein as an indicator, the number of cubic centimeters

10010

of the lime water necessary to neutralize the vinegar, divided by 10, gives the percentage of acetic acid in the vinegar. To make sure that the lime water is saturated, an excess of lime should always be present in the reagent bottle.

Determination of Volatile and Fixed Acids.—Thirty cc. of the vine-gar are transferred to a distilling-flask and subjected to distillation, using a current of steam. Receive the distillate in a 25-cc. graduated cylinder. After 15 cc. have passed over, test from time to time the drops of distillate as they fall into the receiving vessel with litmus-paper, and when free from acid discontinue the distillation. Note the volume of the distillate, mix by shaking, and transfer one-fifth to a white porcelain dish. Titrate as in the case of total acidity, expressing the volatile acids as acetic.

Calculate the fixed acid, expressed in the case of cider vinegar as malic, by subtracting the percentage of volatile acid from the percentage of total acid, and multiplying the result by the factor 1.117. In the case of wine vinegar, express as tartaric acid by using the factor 1.25. To express acidity in terms of sulphuric acid, multiply the percentage of acetic acid by 0.817.

Determination of Alcohol.—Alcohol is present in very small amounts in fruit vinegar that has not been completely acetified. Frear recommends concentrating the distillates as follows: Neutralize 100 cc. of the sample and distill off 40 cc. Then redistill the distillate till 20 cc. have gone over. Cool to 15.6° C. and make up to 20 cc. with distilled water. Determine the specific gravity with a 10-cc. pycnometer, and ascertain from the table on page 661 the per cent by weight of alcohol corresponding to the specific gravity. The percentage in the last distillate, divided by 5, expresses the amount of alcohol in the vinegar.

Detection of Free Mineral Acids.—The ash of genuine cider vinegar is always alkaline. If the ash is neutral, free mineral acids are doubtless present. For their detection the following is a modification of Brannt's method of procedure:

Add to 50 cc. of the vinegar in an Erlenmeyer flask a small bit of starch the size of a wheat-grain, and shake to disseminate it through the fluid. Boil for some minutes, cool, and add a drop of iodine solution. If a blue coloration occurs, no mineral acid is present. In the presence of an appreciable amount of mineral acid, the starch will be converted to dextrin and sugar, and no coloration will be produced by the iodine.

Frear's Method.—Add 5 or 10 cc. of water to 5 cc. of the vinegar, and

to the mixture add a few drops of a solution of methyl violet (one part of methyl violet 2B in 100,000 parts of water). In the presence of mineral acids, a blue or green coloration will be produced.

Determination of Free Mineral Acids.—Hehner's Method.*—To a weighed quantity of the sample add an excess of decinormal alkali, evaporate to dryness, incinerate, and titrate the ash with decinormal acid. The difference between the number of cubic centimeters of alkali added in the first place, and the number of cubic centimeters needed to titrate the ash, represents the equivalent of the free acid present.

Detection and Determination of Sulphuric Acid.—This is determined as barium sulphate by the addition of barium chloride solution. A slight cloudiness on the addition of the reagent indicates the presence of small quantities of sulphate as an impurity, rather than free sulphuric acid. If a minute quantity of free sulphuric acid be present, a rather heavy white cloud on the addition of the barium chloride will be formed, which slowly settles out. According to Brannt, if the quantity of sulphuric acid is more than one part in a thousand, the sulphate of barium formed by addition of the reagent produces a copious precipitate that rapidly falls to the bottom of the receptacle. This may be filtered, washed, ignited, and weighed in the usual manner.

Detection of Free Hydrochloric Acid.—Distill off half of a measured volume of vinegar into the receiving-flask of a distillation apparatus, and to the distillate add a few drops of nitrate of silver reagent. A precipitate indicates hydrochloric acid.

Detection of Malic Acid (Free or Combined).—Absence of malic acid may be assured, if no precipitate occurs with neutral acetate of lead, when a few drops of a solution of this reagent are added to the vinegar. In the presence of malic acid, as in the case of a pure cider vinegar, the precipitate which is formed with lead acetate is flocculent, forms at once, and is of considerable amount. In pure cider vinegar the precipitate will settle to the bottom of the test-tube, leaving a clear supernatant liquid within ten minutes. Unfortunately the acetate of lead test is a negative one, in that several organic acids other than malic will cause a precipitate, as, for instance, tartaric and saccharic acids, the former being found in wine and the latter in molasses vinegar. Malt vinegar also gives a copious precipitate with lead acetate, due to phosphoric acid.

The writer employs the following test † for detecting malic acid in

^{*} Analyst, 1, 1877, p. 105.

[†] An. Rep. Mass. State Board of Health, 1902, p. 485. Food and Drug Reprint, p. 33.

vinegar: Add a few drops of a 10% solution of calcium chloride to some of the vinegar in a test-tube, and make the mixture slightly alkaline with ammonia. Filter off the precipitate that occurs at this point, to the filtrate add two or three volumes of 95% alcohol, and heat to boiling. A copious, flocculent precipitate of calcium malate will form, if malic acid be present, settling to the bottom of the tube in a few minutes. A precipitate will occur in malt and glucose vinegar, due to dextrin.

To confirm the presence of malic acid, filter, wash the precipitate with a little alcohol, dry, dissolve it in strong nitric acid in a porcelain evaporating-dish, and evaporate to dryness over the water-bath, forming calcium oxalate. Boil the residue with sodium carbonate, filter, acidify the filtrate with acetic acid, boil to expel the carbon dioxide, and add a solution of calcium sulphate. A precipitate of calcium oxalate confirms the presence of malic acid.

For the determination of malic acid proceed as directed on page 702. Lead Precipitate.—Hortvet Number.—The quantitative measurement of the precipitate formed with lead acetate, or subacetate, is of considerable importance. Even though the precipitate formed may not be due as was long thought to malic acid, but may be due to phosphoric acid (though this has not been fully proved), it nevertheless remains a fact that the qualitative lead acetate test is one of the most important of all in judging the purity of cider vinegar.

The lead precipitate is best measured as follows: To 25 cc. of the vinegar add 2.5 cc. of U. S. P. subacetate of lead solution. Shake and whirl in a graduated Hortvet tube in the centrifugal machine, and read the volume of the precipitate in the bottom of the tube. The results expressed in cc. on thirty samples of pure cider vinegar are summarized as follows: Highest, 1.4; lowest, 0.5; average, 0.84. The lead number of adulterated cider vinegar runs from a mere trace to 0.5 and sometimes higher.

Winton's Lead Number.—This is determined by the method described for maple products, page 628.

Bailey * obtained by this method the following results:

Cider vinegar (8 samples)	0.075 to 0.290
Malt vinegar (3 samples)	0.158 to 0.548
Distilled vinegar (1 sample)	0.018

^{*}A. O. A. C. Proc., 1908. U. S. Dept. of Agric., Bur. of Chem., Bul. 122, p. 27.

VINEGAR. 769

Hickey* follows the same method, except that he employs only 5 cc. of standard lead subacetate solution and determines the lead in 50 cc. of the filtrate. The lead number found by him in twenty samples of cider vinegar varied from 0.076 to 0.166.

Determination of Acid Tartrate of Potassium.—Berthelot and Fleurien's Method.†—Twenty-five cc. of the vinegar are evaporated on the water-bath to syrupy consistency, and the residue is dissolved in water and made up to its original volume. It is then transferred to a 250-cc. Erlenmever flask, and 100 cc. of a mixture of equal parts of strong alcohol and ether are added, the flask is corked, shaken, and set on ice or in a cold place for forty-eight hours. At the end of this time, if a crystalline precipitate has gathered, the supernatant liquid is decanted upon a filter. and finally the precipitate is washed upon it by a fresh quantity of the ether-alcohol mixture, and the washing continued with this reagent till practically free from acid. The filter and its contents are then transferred to the original flask, and the tartrate is dissolved in boiling water. after which the solution is titrated in the same flask with tenth-normal sodium hydroxide, using phenolphthalein as an indicator. Multiply the number of cubic centimeters of alkali required to neutralize by the factor 0.0188, and the quotient expresses the grams of bitartrate of potash in the sample. Multiply this by 4 to obtain the percentage present.

Polarization and Determination of Sugar.—If the vinegar is light-colored and quite free from turbidity, it may sometimes be polarized undiluted in the 100-mm. tube. Vinegar may often be sufficiently clarified for polarization by filtering twice through the same filter. It is, however, best to add 10% of basic lead acetate solution, and to filter before polarizing, thus removing the malic or tartaric acids which may have a slight effect on the polarization. In case of dark-colored or turbid samples, add to 50 cc. of the sample 5 cc. of about equal quantities of lead subacetate and alumina cream, shake, filter, and polarize in a 200-mm. tube, adding 10% to the reading on account of the dilution. The polarization value of the vinegar is conveniently expressed in terms of actual direct reading obtained by the undiluted sample in a 200- or 400-mm. tube.

If the invert reading is desired for calculation of sucrose or commercial glucose, subject the sample to inversion with hydrochloric acid and heat, as in the case of sugars.

^{*} Ibid.

[†] Girard et Dupré, Analyse des Matières Alimentaires, p. 128

For the determination of sucrose, use Clerget's formula (p. 588), calculating the true direct and invert readings from the direct and invert readings of the undiluted vinegar on the basis of the normal weight of the sample, by multiplying the obtained readings by 0.26 in the case of the Soleil-Ventzke instrument.

Determination of Reducing Sugars before and after Inversion.—Two portions of 25 cc. each are measured into 100-cc. flasks. One portion is diluted with 25 cc. of water, 5 cc. of concentrated hydrochloric acid are added, and the solution subjected to inversion by heating to 70° for 10 minutes and cooling. Both portions are neutralized with sodium hydroxide and made up to the mark. The reducing sugars are determined in each portion by Defren's modification of O'Sullivan's method (page 594) and calculated as dextrose.

Levulose may be determined as on page 626, polarizing the vinegar at two different temperatures.

ADULTERATION OF VINEGAR.

Standards of Purity.—In nearly all localities where pure-food laws prevail there are special provisions setting forth the requirements of pure vinegar as to percentage of acids, solids, and other conditions, differing considerably with the character of the vinegar used. Thus, in England, where the principal vinegar is malt vinegar, the legal standards are considerably different from those in force in France and Germany, where wine vinegar is prevalent. These differ again from the requirements found in the United States and Canada, where cider vinegar is the chief product.

Most of the state food laws fix a standard for the acidity of cider vinegar varying from 3.5 to 4.5 per cent of acetic acid, and in most cases also a minimum standard for total solids or residue of from 1.5 to 2 per cent. Special laws stipulate furthermore in some states that cider vinegar, sold as such, must be exclusively the product of pure apple cider. In such cases cider vinegar may be adulterated by non-conformance to the standard in either acidity or solids or both, while yet it may be exclusively made from pure apple cider. This may be due either to actual watering or to incomplete acetification. On the other hand, so-called cider vinegar may be of legal standard as to solids and acidity, and yet be entirely spurious.

Following are the U. S. standards for the various vinegars:

Vinegar, Cider Vinegar, Apple Vinegar, is the product made by the alcoholic and subsequent acetous fermentations of the juice of apples, is lævo-rotatory, and contains not less than 4 grams of acetic acid, not less than 1.6 grams of apple solids, of which not more than 50% are reducing sugars, and not less than 0.25 gram of apple ash in 100 cc. (20° C.); and the water-soluble ash from 100 cc. (20° C.) of the vinegar contains not less than 10 milligrams of phosphoric acid (P_2O_5) , and requires not less than 30 cc. of decinormal acid to neutralize its alkalinity.

Wine Vinegar, Grape Vinegar, is the product made by the alcoholic and subsequent acetous fermentations of the juice of grapes, and contains in 100 cc. (20° C.), not less than 4 grams of acetic acid, not less than 1.0 gram of grape solids, and not less than 0.13 gram of grape ash.

Malt Vinegar is the product made by the alcoholic and subsequent acetous fermentations, without distillation, of an infusion of barley malt, or cereals whose starch has been converted by malt, is dextro-rotatory, and contains, in 100 cc. (20° C), not less than 4 grams of acetic acid, not less than 2 grams of solids, and not less than 0.2 gram of ash; and the water-soluble ash from 100 cc. (20° C), of the vinegar contains not less than 9 milligrams of phosphoric acid (P₂O₅), and requires not less than 4 cc. of decinormal acid to neutralize its alkalinity.

Sugar Vinegar is the product made by the alcoholic and subsequent acetous fermentations of solutions of sugar, syrup, molasses, or refiners' syrup, and contains, in 100 cc. (20° C), not less than 4 grams of acetic acid.

Glucose Vinegar is the product made by the alcoholic and subsequent acetous fermentations of solutions of starch sugar or glucose, is dextrorotatory, and contains, in 100 cc (20° C.), not less than 4 grams of acetic acid.

Spirit Vinegar, Distilled Vinegar, Grain Vinegar, is the product made by the acetous fermentation of dilute distilled alcohol, and contains, in 100 cc. (20° C.), not less than 4 grams of acetic acid.

Accidental Adulteration of vinegar may result in the presence of injurious metallic salts, such as of copper, lead, or zinc, derived from vessels or utensils used in the manufacture of vinegar, or even minute traces of arsenic may be found, when glucose has been employed as an ingredient

or source of the vinegar, the arsenic being in this case probably due to impure sulphuric acid used in the manufacture of the glucose.

Willful or Fraudulent Adulteration is, however, common, in which misbranded vinegar is sold under names suggesting a class other than that to which it really belongs, or wherein entirely artificial substitutes are made up for pure cider, malt, or wine vinegar, in which the color, residue, and acid principle may be either or all of spurious origin.

Artificial Cider Vinegar is in most cases readily detected, though very ingenious imitations are on the market, involving not a little skill and chemical knowledge in their manufacture.

Entirely artificial substitutes for cider vinegar are frequently made up of spirit vinegar, colored with caramel, and having the solids reinforced by apple jelly, made for the most part out of exhausted apple pomace, which is the residue left after the apple-stock has been subjected to one and sometimes two pressings. The jelly used for this purpose is not infrequently made up with commercial glucose. All grades of adulterated vinegar are to be found, from the wholly spurious substitute above described, to the varieties in which cider vinegar is itself present, but is pieced out or reinforced by the admixture of coloring matter, mineral acid, wood vinegar, or of molasses or glucose vinegar. Acetic ether is sometimes employed to impart flavor to the product. All the characteristics of a pure cider vinegar are difficult to duplicate artificially, though some of them may be.

Character of the Residue.—The residue of pure cider vinegar should be thick, light brown in color, of a viscid or mucilaginous consistency, somewhat foamy, having an astringent acid though pleasant taste very suggestive of baked apples, which it also resembles in odor. The odor of molasses is very apparent in the residue of vinegar having sugar-house wastes, and the smell of a malt-vinegar residue is also very characteristic. If pyroligneous or wood vinegar has been introduced, the dried residue will have a tarry or smoky taste and smell.

The residue of cider vinegar is very soluble in alcohol, while that of malt vinegar is only slightly soluble. Wine vinegar residues dissolve readily in alcohol, except for the granular residue of cream of tartar. If the loop of a clean platinum wire be rubbed in the vinegar residue and ignited in a colorless Bunsen flame, the color imparted will, if the vinegar has been made from pure cider exclusively, consist altogether of the palelilac color of a potash salt without any of the yellow sodium flame being

VINEGAR. 773

visible. In all vinegars other than of pure cider, the sodium flame will predominate, when the residue is burnt as above. Again, the ignited residue left in the loop of wire in the case of a pure cider vinegar will form a fusible bead, having a strong alkaline reaction upon moistened test-paper, and effervescing briskly when immersed in acid. The presence in vinegar of even a slight trace of added mineral acid will prevent the ignited residue from having the alkaline reaction, or effervescing with acid.*

The residue of malt or beer vinegar is brown and gummy, containing a considerable quantity of dextrin. Not only are the appearance and odor of the dried vinegar residue to be particularly noted, but also the odor given off in the first stages of burning this residue to an ash. With cider vinegar the apple odor is very marked while burning. In vinegar wherein molasses products have been employed, the smell of charred sugar is usually apparent, while with glucose vinegar the smell of burnt corn predominates.

On burning the residue of malt vinegar, the odor produced at first is not unlike that of toasted bread. At a later stage in the burning the vapors evolved are very pungent.

The Character of the Ash is of considerable importance in determining the source of a sample of vinegar. The ash of pure cider and malt vinegar is quite strongly alkaline, while that of distilled and wood vinegar is only slightly alkaline. The ash of cider vinegar is high in alkaline carbonates.

In cider and malt vinegar the quantity of phosphoric acid present in the ash is considerable, while only traces are present in distilled or spirit vinegar. Considerably more than half the phosphoric acid in the ash of cider vinegar is soluble, while no soluble phosphoric acid is present in the ash of spirit vinegar.

The percentage of ash in total solids is of some value in judging the purity of cider vinegar. According to Frear.† if the ash of the vinegar is less than 10% of the total solids, the vinegar may be suspected of having added unfermented material, while a percentage of ash less than 6 is absolute evidence that the vinegar is not genuine cider vinegar.

The alkalinity of 1 gram of the ash of pure cider vinegar should be

^{*} Davenport, 18th An. Rep., Mass. Board of Health, 1887, p. 150.

[†] Report of Penn. Dept. of Agric., 1898, p 38.

equivalent to at least 65 cc. of tenth-normal acid. At least 50% of the phosphates in the ash should be soluble in water.

Character of the Sugars.—One of the most important steps in establishing the source of a vinegar consists in subjecting it to polarization (p. 769). From the nature of the sugar-content of the apple juice, not only when freshly expressed, but also when allowed to undergo alcoholic fermentation, and, furthermore, after it has gone over into vinegar, the polarization through all three stages is always left-handed.

Browne * has shown that the optical rotation of the freshly expressed juice of eleven varieties of apple varies from 19.24° to 49° to the left on the Ventzke scale, in a 400-mm. tube. Also that in the case of five samples of completely fermented cider, examined five or six months after pressing, the left-handed rotation in a 400-mm. tube varied from 1.76° to 5.28°. He showed, furthermore, that a sample of pure cider jelly made up of concentrated apple juice had a left-handed rotation amounting to 21.35° in a 200-mm. tube (20 grams made up 100 cc.), and finally that four cider vinegar samples of known purity showed left-handed readings of from 0.96° to 2.94° Ventzke in a 400-mm. tube.

The feft-handed rotation of pure cider vinegar is a characteristic so fixed and unalterable that a right-handed polarization of more than 0.5° may safely be assumed as evidence of adulteration. The polarization of cider vinegar, expressed in terms of 200 mm. of the undiluted sample should lie between -0.1° and -4.0° Ventzke. If the direct polarization of a sample of vinegar is right-handed, while the invert is left-handed, sugar-house wastes or molasses may be suspected as an adulterant.

If both direct and invert readings are right-handed, commercial glucose is undoubtedly present. If the polarization of the vinegar is far to the left, unfermented cider jelly has probably been used to reinforce the solids.

Frear regards the ratio of reducing sugars after inversion to total solids as a useful factor in discriminating between pure cider vinegar and the common artificial substitutes in which the solids of distilled vinegar are reinforced by apple jelly, or in which commercial glucose or molasses vinegars are used. When the reducing sugars after inversion form more than 25% of the entire solids, the alleged cider vinegar is undoubtedly

^{*} Bull. 58, Penn. Dept. of Agric., "A Chemical Study of the Apple and Its Products."

spurious. In pure cider vinegar the per cent of reducing sugar is the same after inversion as before. The same is true of glucose vinegar

Vinegar containing added molasses or cane sugar will, however, naturally show an increase in reducing sugar after inversion.

A large content of alcohol in cider vinegar, otherwise showing the constants of pure vinegar except for the low acidity, would indicate incomplete acctification. A high content of nitrogen is characteristic of malt vinegar.

Data of analyses of samples of vinegar examined in the Food and Drug Department of the Massachusetts State Board of Health are given in the tables on this page and the next. The table below shows in summarized form the results obtained from the examination of eighty-four samples of undoubtedly pure cider vinegar examined in 1901.*

CIDED	VINECAD	FOUND	DIIDE

	Acid (Per Cent).	Solids (Per Cent).	Ash (Per Cent).	Polarization.
MaximumMinimum.		4.00	0.58	-5-4 -0.4
Mean		2.43	0.38	-2.0

The second table includes samples of adulterated vinegar, sold for cider vinegar, none of which were probably made from cider. It will be noticed that in several of the samples the amount of glucose was abnormally large, as is shown by the very high right-handed polarization, in one case amounting to over 12°.

Direct Tests Made on the Vinegar.—The genuine or spurious nature of cider vinegar may usually be established by direct tests with reagents on the vinegar itself. The appearance, taste, and odor of the vinegar should be noted. Brannt † applies the test of odor in vinegar as determining its character, by rising out a large beaker with the sample, and

^{* 32}d An. Rep. (1900), p. 661, Food and Drug Reprint, p. 44; 33d An. Rep. (1901), p. 467, Food and Drug Reprint, p. 47; 34th An. Rep. (1902), p. 483, Food and Drug Reprint, p. 31.

[†] A Practical Treatise on the Manufacture of Vinegar, p. 219.

WINECAD	NOT	THE	EVCI HEIVE	PRODUCT OF	DIIDE	ADDIE CI)ED
VINCOAR	NOI	Inc	LACLUSIVE.	PRODUCT OF	PURE	APPLE UII	Jr.K.

Per Cent Acetic Acid.	Per Cent Total Solids.	Per Cent Ash.	Per Cent Ash in Total Solids.	Polarization in 200-mm. Tube.	Lead Acetate.
5.90	-40	••••		+1.4	No precipitate
5.14	.36	••••	••••	.0	" "
5.12	-53	••••		+ .6	1
4.83	3.70	.32	8.65	+8.0‡	1
4.82	2.71	-13	4.80	+9.6	Heavy precipitate*
4.80	1.97	.20	10.15	+ .9	Precipitate
4.80	1.03	-27	14-75	+1.1	"
4.66	2.92	.20	6.49	+2.2	No precipitate
4.60	2.57			+2.6	11 11
4.56	2.60			+7.0‡	†
4-54	3-97	.19	4.78	+5.6	No precipitate
4.54	3.90	- 32	9.72	+5.0	" "
4-54	2.94	.23	7.82	+5.0	"
4.54	2.70	.23	8.52	+ -4	Precipitate
4.50	3.05			+2.2	No precipitate
4.50	2.92	.22	7.52	+ .9	" "
4.50	2.69	• • • •		+2.8	"
4.48	3.80			+ 12.0‡	"
4.46	2.80			+2.6	" "
4.42	2.75			+3.2	Slight precipitate
4.42	2.10			+9.2	Precipitate
4.40	2.51	.20	11.15	+1.1	11
4.40	-97			+ .4	No precipitate
4.38	.29			+1.6	· · · · · · · · · · · · · · · · · · ·
4 - 32	.70	.09	12.86		"
4.08	3.35	••••	l	+1.2	Precipitate
3.98	-55			+1.8	Slight precipitate

^{*} Cider vinegar to which apple jelly containing glucose had been added for the purpose of increasing the solids after watering.

† This sample contained a large amount of phosphate, and consequently the test for malates is obscured.

† These samples polarized practically the same after as before inversion, indicating much glucose.

after allowing it to stand for some hours, examining the few drops remaining in the beaker. The acetic acid having for the most part become volatilized, the characteristic vinous odor of pure wine vinegar would at this stage be very prominent, while that of cider vinegar would be entirely different. The odor of the two vinegars is very similar in their ordinary state. The peculiar fruity flavor of pure cider-vinegar is very characteristic and not readily imitated by cheaper substitutes. Only a very slight turbidity should be produced in pure cider vinegar by the addition of either ammonium oxalate (absence of lime), barium chloride (absence of sulphuric acid or sulphate), and nitrate of silver (absence of hydrochloric acid or chlorides).

Vinegar in which glucose has been used nearly always gives a precipitate with ammonium oxalate, due to the sulphate of calcium present.

VINEGAR. 777

The character of the precipitate produced by neutral lead acetate should be particularly noted. Unless it is flocculent and copious, settling out after a few minutes, cider vinegar is not pure, even if a marked turbidity is produced. Added apple jelly from exhausted apple pomace gives such a turbidity, and is to be suspected when not more than a cloudiness is produced on addition of the lead acetate reagent. Pure cider vinegar should respond in a perfectly normal manner to both the lead acetate and the calcium chloride tests for malic acid.

Wood Vinegar or Pyroligneous Acid is sometimes rendered apparent by the empyreumatic or tarry taste and odor imparted to the product. When, however, the added acetic acid has been so purified that the tarry taste and odor are lacking, its presence may often be proved by the traces of furfurol which always accompany it.

Test for Furfural.—A little of the vinegar is subjected to distillation, and to the first few drops of the distillate is added a little colorless anilin solution. A fading crimson color will be produced in presence of furfurol. This reaction may sometimes be obtained upon the vinegar itself without distillation, if sufficient added wood vinegar be present.

The first portion of the distillate of wood vinegar reduces permanganate of potassium to a marked degree.

The Addition of Spices to vinegar in order to increase the pungency is best detected by first neutralizing the vinegar with sodium carbonate and then tasting. Under these conditions, the admixture of spices is rendered very apparent.

Detection of Caramel.—Considerable added caramel in vinegar is apparent from the unnaturally dark color and extremely bitter taste of the residue after evaporation.

Tests for caramel made on the vinegar residue, if long dried at the temperature of the water-bath, are not to be depended on as establishing the presence of added caramel, since at that temperature the decomposition of the sugar may in any event cause a positive test.

Caramel is detected by Crampton and Simon's and Amthor's tests (p. 752). A further indication of caramel is the reducing power of the water solution of the precipitate obtained in Amthor's test.

Examination for Metallic Impurities.—Lead and Zinc are best looked for in the ash of the vinegar in cases where, like cider vinegar, the percentage of extract is high. A large volume of the vinegar is evaporated to substantial dryness over the water-bath. This may most readily be done in a 100-cc. platinum wine-shell, adding the vinegar in successive

portions. To the residue add a small amount of sodium hydroxide, and burn to an ash in a muffle, or over a low flame, using potassium nitrate if necessary, a little at a time. Take up the ash in dilute hydrochloric acid, and examine for lead and zinc as in the case of canned goods.

In the case of vinegar low in extract, as in spirit vinegar, the sample may be evaporated to dryness, the residue dissolved directly in dilute hydrochloric acid without ignition, and the acid solution subjected to direct examination for lead and zinc.

Copper is best determined by electrolysis. 100 cc. of the vinegar are evaporated to a volume of about 10 cc. with a little sulphuric acid, filtered into a platinum dish, and subjected to electrolysis, using conveniently the apparatus described on page 608.

Arsenic.—Boil down a portion of the vinegar, to which concentrated nitric acid has been added, to a small volume, then add a few cubic centimeters of concentrated sulphuric acid, and continue the heating till fumes of sulphuric acid show the nitric to have been driven off. Cool, dilute with water, and test in the Marsh apparatus.

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CHAPTER XVII.

ARTIFICIAL FOOD COLORS.

THE use of artificial dyestuffs in food products has greatly increased during recent years, both in degree and in variety of colors employed. Where formerly but a few well-known coloring matters, chiefly so-called vegetable colors and occasionally mineral pigments were used for this purpose, a vast array of dyes, chosen largely from the coal-tar colors, are now found in food, so that at present the exact identification of the particular dyestuff employed in all cases presents a somewhat formidable problem to the analyst. The problem may consist in determining the class to which a commercial food color or combination of colors belongs, or it may consist in isolating the color itself, and afterwards identifying it as far as possible, for the purpose of determining whether or not it is harmless within the meaning of the law.

The effect of imparting to the cheaper varieties of jellies, jams, and ketchups which flood the market such intense and striking colors that these products in no wise resemble their pure uncolored prototypes, has a tendency in many cases to mislead the public into the idea that the genuine products are inferior by contrast, and to create a craving for unnaturally colored varieties. Indeed, the adherents to the free use of coloring matters in food assert that these brilliant hues please the eye and are hence legitimate.

Objectionable Features.—With the exception of confectionery and certain dessert preparations, in which dyes may be employed purely for æsthetic considerations only (a fact which is well understood by the consumer), the use of coloring matters in food is mainly for the purpose of deceiving as to their true character. The use of dyestuffs in food is objectionable on two accounts, first as introducing in some cases materials injurious to health, and second, in nearly all cases as deceiving the purchaser by concealing inferiority, or by making the goods

780

appear of greater value than they really are. In most states the food laws regarding employment of colors are so framed, that the presence of such colors constitutes an offense under one or the other of the above heads, mainly, however, because, by reason of their use, cheaper or inferior materials are made to masquerade for the higher or genuine grades, as, for instance, when alleged currant jelly is found to consist chiefly of apple-stock and commercial glucose, colored with an artificial red dye.

In such cases the analyst has merely to prove conclusively that an artificial color is present, even if he does not identify the dye itself. It is of course more satisfactory to at least show in addition whether the dye present is of vegetable origin, or is of the coal-tar variety, and in most cases this can readily be done, even if it is not easy to identify the exact color.

In localities where laws prevail stipulating that what are commonly known as "mixtures" or "compounds" to be legally sold, must be labeled with the names and percentages of ingredients, the law applies to coloring matters as well as other ingredients, and the exact dye or dyes employed should appear on the label. Otherwise the product must be classed as adulterated.

Toxic Effects of Colors.—Formerly the use of such pigments as chromate of lead was common in coloring confectionery, but lead chromate is rarely used at present. Other mineral pigments obviously unfit for use in food by reason of their well-known poisonous effects are those which contain salts of arsenic, mercury, lead, and copper. While most of the coal-tar colors are considered harmless in themselves, some are decidedly objectionable, and should not be used in foods. Under the latter class are included, first, those in connection with the manufacture of which arsenic, mercury, or other poisonous mineral ingredients have been used, such for example as arsenical fuchsin, and, second, those which are themselves inherently poisonous, as for instance picric acid. Fuchsin is now largely made without the aid of arsenic acid, and this variety is, perhaps, harmless. The toxic effects of many of the coal-tar colors have not been thoroughly established excepting in a negative way. Weyl has made many experiments on dogs and rabbits in which these animals have been fed with varying amounts of coloring material. In nearly all cases the doses far exceeded the amounts ordinarily taken in food, and the experiments are of value mainly in so far as they show harmless results of certain colors on the animal. It is to be regretted

that physiological experiments cannot more readily be tried on human beings, so as to study the effects of administering to them such amounts as are used in food.

More conclusive results (though still of a negative character) tending to establish the harmlessness of most of the coal-tar colors are given by Grandhomme * in statistics showing the condition of health of laborers in factories where these dyestuffs are made, in comparison with those engaged in other industries where poisonous materials are handled. From these it appears that the proportion of illness among the anilinmakers is remarkably small.

In the case of coloring confectionery by the use of mineral pigments, a considerable amount of the coloring material must be used, forming without doubt a source of danger in some cases. With coal-tar dyes, on the contrary, the case is different. One ounce of auramine, for instance, has been found sufficient to give a deep-yellow color to 2,000 pounds of confectionery, so that almost an infinitesimal amount of the actual dyestuff is taken into the system. Hence it is that very little danger need be apprehended from the use of most coal-tar colors in food, objectionable as they certainly are as a commercial fraud.

Injurious and Non-injurious Colors.—Various countries have enacted specific laws regulating the use of coloring matters in foods, especially England, France, Germany, Austria, and Italy. In some cases attempts have been made to specify harmful and harmless colors. The National Confectioners' Association of the United States has compiled a useful classified list of injurious and harmless colors,† the classification being based largely on the results of experiments by Weyl and König, as well as upon the Resolutions of the Association of Swiss Chemists, and on the French Ordinances regarding food colors. The list is as follows:

HARMFUL MINERAL COLORS.

Compounds of Copper.—Blue ashes, mountain blue, etc.

Compounds of Lead.—Massicot, red lead, white lead, Cassel yellow, Paris yellow, Turner yellow, Naples yellow, sulphates of lead, chrome yellow, Cologne yellow, etc.

Compounds of Barium.—Ultramarine yellow, etc.

^{*} Weyl, Sanitary Relations of the Coal-tar Colors, pp. 28-30.

[†] Colors in Confectionery. An Official Circular from the Executive Committee of the National Confectioners' Association of the U. S., 1899.

Compounds of Mercury.—Vermilion, etc.

Compounds of Arsenic.—Scheele's green, Schweinfurth green, etc.

In Other Words colors in whose preparation mercury, lead, copper, arsenic, antimony, tin, zinc, chromium, and barium compounds are used.

HARMFUL ORGANIC COLORS.

Red Colors.—Ponceau 3RB.—Ponceau B extra, fast ponceau B, new red L, scarlet EC, imperial scarlet, old scarlet, Biebrich scarlet.

Crocein Scarlet 3B.—Ponceau 4RB.

Cochenille Red A.—Crocein scarlet 4B and G, brilliant scarlet, brilliant ponceau 4R, ponceau 4R, ponceau brilliant 4R, new coccin, scarlet.

Crocein Scarlet 7B.—Crocein scarlet 8B, ponceau 6RB.

Crocein scarlet O extra.

Safranin.—Safranin T, safranin extra G, safranin G extra GGSS, safranin GOOO, safranin FF extra No. O, safranin conc., safranin AG extra, safranin AGT extra, anilin pink.

Yellow Colors.—Gum gutta.

Picric acid.

Martius Yellow.—Naphthylamin yellow, jaune d'or, Manchester yellow, naphthalin yellow, naphthol yellow, jaune naphthol.

Acme Yellow.—Chrysoin, chryseolin yellow T, gold yellow, resorcin yellow, acid yellow RS, tropæolin O, jaune II.

Victoria Yellow.—Victoria orange, anilin orange, dinitrocresol, saffron substitute, golden yellow.

Orange II.—Orange No. 2, orange P, orange extra, orange A, orange G, acid orange, gold orange, mandarin G extra, beta-naphtholorange, tropæolin OOO No. 2, mandarin, chrysaurin.

Metanil Yellow.—Orange MN, tropæolin G, Victoria yellow (O double conc.), jaune G (metanil extra).

Sudan I.—Carminaph.

Orange IV.—Orange No. 4, orange N, orange GS, new yellow, acid yellow D, tropæolin OO, fast yellow, diphenylorange, diphenylamine orange, jaune d'anilin, anilin yellow.

Green Colors.—Naphthol green B.

Blue Colors.—Methylene blue BBG.—Methylene blue BB, in powder extra, methylene blue DBB extra, methylene blue BB (crystalline) ethylene blue.

Brown Colors.—Bismarck Brown.—Bismarck brown G, Manchester brown, phenylen brown, vesuvin, anilin brown, leather brown, cinnamon brown, canelle, English brown, gold brown.

Vesuvin B.—Manchester brown EE, Manchester brown PS, Bismarck brown, Bismarck brown T, brun Bismarck EE.

Fast Brown G.—Acid brown.

Chrysoidin.—Chrysoidin G, chrysoidin R, chrysoidin J, chrysoidin Y.

HARMLESS MINERAL COLORS.

Blue Colors.—Ultramarine blue.

Violet Colors.—Ultramarine violet.

Brown Colors,-Manganese brown.

Chocolate-brown and colors of a similar nature have as their basis natural or precipitated oxide of iron, which in an impure condition may have small quantities of arsenic in its composition. It is possible with proper care to secure a raw material entirely free from this objectionable element, and no oxide of iron containing any traces of arsenic should be used in the preparation of color.

Green Colors.—Ultramarine green.

HARMLESS ORGANIC COLORS.

Red Colors.—Cochineal carmine.

Carthamic acid (from saffron).

Redwood.

Artificial atizarin and purpurin.

Cherry and beet juices.

Eosin.—Eosin A, eosin G extra, eosin GGF, eosin water soluble, eosin 3J, eosin 4J extra, eosin extra, eosin KS ord., eosin DH, eosin JJF.

Erythrosin.—Erythrosin D, erythrosin B, pyrosin B, primrose soluble, eosin bluish, cosin J, dianthin B.

Rose Bengale.—Rose bengale N, Rose bengale AT, rose bengale G, bengalrosa.

Phloxin.—Phloxin TA, eosin blue, cyanosin, eosin 10B.

Bordeaux and Ponceau reds, resulting from the action of naphthol-sulphonic acids on diazoxylene:

Ponceau 2R.—Ponceau G, ponceau GR.

Ponceau R.—Brilliant ponceau G, ponceau J.

Bordeaux B.-Fast red B, Bordeaux R extra.

Cerasin.-Rouge B.

Ponceau 2G.—Brilliant ponceau GG, ponceau JJ.

Fuchsin S.—Acid magenta, rubin S, fuchsin acide (free from arsenic). Archil Substitute.—Naphthion red.

Orange I.—Orange No. 1, naphtholorange, alpha-naphtholorange, tropæolin OOO No. 1.

Congo red.

Azorubin S.—Azorubin, azorubin A, azoacidrubin, fast red C, carmoisin, brilliant carmoisin O, rouge rubin A.

Fast Red D.—Fast red EB, fast red NS, amaranth, azoacidrubin 2B, Bordeaux DH, Bordeaux S, naphthol red S, naphthol red O, Victoria ruby, wool red (extra), cenanthinin.

Fast Red.—Fast red E, fast red S, acid carmoisin S.

Ponceau 4GB.—Crocein orange, brilliant orange G, orange GRX, pyrotin orange, orange ENL.

Fuchsin.

Metanitrazotin.

Yellow and Orange Colors.—Annatto.

Saffron.

Safflower.

Turmeric.

Naphthol Yellow S.—Citronin A, sulphur yellow S, jaune acide, jaune acide C, anilin yellow, succinine, saffron-yellow, solid yellow, acid yellow S.

Brilliant Yellow .-- (Sch.)

Ponceau 4GB. — Crocein orange, brilliant orange G, orange GRX, pyrotin orange, orange ENL.

Fast Yellow.—Fast yellow G, fast yellow (greenish), fast yellow S, acid yellow, new yellow L.

Fast Yellow R.—Fast yellow, yellow W.

Azarin S.

Orange I.—Orange No. 1, naphtholorange, alpha-naphtholorange, tropæolin OOO No. 1.

Orange.—Orange GT, orange RN, brilliant orange O, orange N.

Mixtures of harmless red and yellow colors.

Green Colors.—Spinach green.

Chinese green.

Malachite Green.-Malachite green B, benzaldehyde green, new Vic-

toria green, new green, solid green crystals, solid green O, diamond green, bitter amond oil green, fast green.

Dinitrosoresorcin.—Solid green O in paste, dark green, chlorine green, Russia green, Alsace green, fast green, resorcinol green.

Mixtures of harmless blue and yellow colors.

Blue Colors.—Indigo.

Litmus.

Archil blue.

Gentian Blue 6B.—Spirit blue, spirit blue FCS, opal blue, blue lumière, Hessian blue, light blue.

Coupiers Blue.—Fast blue R and B, solid blue RR and B, indigin DF, indulin (soluble in alcohol), indophenin extra, blue CB (soluble in alcohol), nigrosin (soluble in alcohol), noir CNN.

In General such blues as are derived from triphenylrosanilin or from diphenylamin.

Violet Colors. — Paris Violet. — Methyl violet B and 2B, methyl violet V3, pyoktanin coeruleum, malbery blue.

Wool black.

Naphthol black P.

Azoblue.

Mauvein.—Rosolan, violet paste, chrome violet, anilin violet, anilin purple, Perkins violet, indisin, phenamin, purpurin, tyralin, tyrian purple, lydin.

Brown Colors.—Caramel.

Licorice.

Chrysamin R.

Use of Colors in Confectionery.—Regarding the choice of colors for use in confectionery and precautions to be observed in their use, the Confectioners' Association has offered the following considerations:

First. That coal-tar colors are specially adapted to the wants of confectioners on account of their brilliancy, permanency, and high coloring power, by reason of which last-named quality only infinitesimal amounts of color need be or can be used to give the desired effects.

Second. That there is no evidence to show that any poisonous or hurtful colorings have in recent years been found in confectionery. Reports of deaths from poisoned candy are only too frequently made, but no autopsy has ever been published confirming them.

Third. That while the exceedingly small proportions of color used in confectionery constitute a practical safeguard to the public health, con-

fectioners are in duty bound to provide against all possible contingencies of harm, by using the utmost care in obtaining absolutely non-poisonous colors, buying only from color-dealers of established reputation and unquestioned responsibility, whose colors are tested at frequent intervals, and are vouched for by competent chemists.

Confectioners should require that a guarantee be put upon each package of color, stating not only that the contents are non-poisonous, but also that they will not in any way interfere with digestion or injure health.

Fourth. Any illegitimate use of coloring matter in confectionery as a substitute for chocolate or any other material or ingredient, or for the purpose of adding bulk or increasing the weight of the confectionery in which it is incorporated, should not be permitted or countenanced. Both the letter and the spirit of these laws should clearly prevent the illegitimate use of coal-tar colors or of earth colors, such, for example, as chocolate-brown, coconole brown, or chocolatina.

Fijth. That color-dealers furnishing colors to confectioners should publish printed lists of their colors under the various names and titles by which they are known and offered for sale, accompanying such lists with ample certifications by competent chemists to their purity and suitableness for coloring confectionery and other articles of food. They should also attach to each package or other container of color a guarantee that it does not contain anything injurious to health.

VEGETABLE COLORS.

These with a few mineral pigments and cochineal were formerly almost exclusively used for coloring food products, and are still used to some extent.

Most of the vegetable colors, according to L. Robin,* react with ammonia to form a coloration, usually passing from violet to blue, then to a brownish green, when the ammonia is added little by little in excess to the color in solution. If by the addition of ammonia to a solution of an unknown color the green coloration does not result, the presence may be suspected of orchil or cudbear, logwood, cochineal, or a coal-tar dye.

The following vegetable colors are occasionally found in food, with some of the reactions in aqueous solution, as given by Robin:

^{*} Girard et Dupré, Analyse des Matières Alimentaires, p. 579.

RED COLORS.

Nature of Color.	Ammonia.	Alum and Sodium Carbonate 20% Solution. Mixture of Aluminum Acetate and Acetate acetate and Acetate acetate and Acetate aceta			
		Lake.	Filtrate from	Sodium Carbonate.	
Bilberry (whor-tleberry)	Dull greenish	Greenish blue, rose-colored on edges		Bluish violet	
Beet	Muddy yellow, brown, or rose- red			Garnet	
Black currant	Deep green	Greenish blue	Bottle-green	Violet-blue	
Logwood	Red tinged with violet	Blue tinged with violet		Tinged with violet	
Brazil wood	Currant-red	Rose	Rose tinged	Lilac to wine color	
Raspberry	Bluish green	Rose tinged	Bluish green	Lilac tinged with violet	
Currant	Yellow-brown to greenish	Gray to lilac	Dull maroon to bottle-green	Red-maroon	
Blackberry	Yellowish green	White or rose vio-	Bluish	Dull violet	
Phytolacca	Lilac	Violet		Clear violet, pass- ing to yellow with ammonia	
Elderberry	Light green	Blue tinged with violet		Violet, quickly passing to blue with acetate of copper	

YELLOW COLORS.

Nature of Color.	Ammonia.	Hydrochloric Acid.	Alum and Carbonate of Soda 20% Solution. Lake.
Persian berries	Yellow-red	Precipitate yellow- brown	Orange
Old fustic	Very bright yellow Becomes clearer	Yellow-orange Bright yellow pre- cipitate	Orange Yellow-red tending to green
Young fustic Turmeric	Yellowish red Brown-red	Becomes yellower Crimson precipitate	Bright yellow Bright yellow

Additional yellow vegetable colors sometimes used in foods are the following, taken from a table of Leed's,* showing reactions given by treating a few drops of an alcoholic solution of the color with an equal volume of the reagent.

Most of these vegetable colors do not directly dye wool or silk a fast color, but as a rule require the use of a mordant. Many of these colors may be fixed on cotton (previously mordanted by boiling in a solution

REACTIONS	OF	COLOBING	MATTEDS
REACTIONS	UF	COLORING	MALIERS.

Coloring Matter.	Concentrated H ₂ SO ₄ .	Concentrated HNO ₈ .	H ₂ SO ₄ +HNO ₃ .	Concentrated HCl.
Annatto	Indigo-blue, chang- ing to violet	Blue, becoming colorless on standing	Same	No change, or only slight dirty yel- low and brown
Turmeric	Pure violet	Violet	Violet	Violet, changing to original color on evaporation of HCl
Saffron	Violet to cobalt blue, changing to reddish brown	Light blue, chang- ing to light red- dish brown	Same	Yellow, changing to dirty yellow
Carrot	Umber brown	Decolorized	Same with NO ₂ fumes and odor of burnt sugar	No change
Marigold	Dark olive-green, permanent	Blue, changing instantly to dirty yellow-green	. Green	Green to yellowish green
Safflower	Light brown	Partially decolor- ized	Decolorized	No change

of aluminum acetate or potassium bichromate) by boiling the mordanted fibers in a bath of the colored solution, rendered acid by acetic acid. The dyed fibers are then examined by reagents, as in tables given on pages 804-11.

SPECIAL TESTS FOR VEGETABLE COLORS.—Orchil and Cudbear, both derived from lichens, dye wool red in acid bath. The colored fiber, in the case of cudbear, is turned blue by treatment with ammonia. For reactions of orchil on the fiber see table, page 807. Robin's test for orchil in aqueous solution consists in shaking it with ether, which, if orchil is present, is colored yellow. On treatment of the ether with ammonia, the yellow color is changed to blue, and, by adding acetic acid, goes over to a reddish violet.

Logwood, according to Robin, in aqueous solution colors ether yellow, and on treating the ether with ammonia the color becomes red or faintly violet. Potassium bichromate gives a violet coloration, mingled with greenish yellow. If cotton is first mordanted by boiling with aluminum acetate, it is dyed violet when boiled in a solution of logwood.

Turmeric is best extracted from a dry residue with alcohol, which it colors yellow. The color is transferred to a piece of filter-paper by soaking the paper in the alcoholic tincture, the paper is dried and dipped in a dilute solution of boric acid or borax slightly acidulated with hydrochloric acid. On again drying the paper, it will be of a cherry-red color if turmeric is present, and when touched with a drop of dilute alkali will turn dark olive.

Caramel.—Care should be taken in testing for caramel not to subject the sample to long-continued heating, even on the water-bath. Indeed caramel is sometimes developed spontaneously in saccharine food products during their process of manufacture when heat is used, by the charring of the sugar. If solutions are to be concentrated or brought to dryness before testing for caramel, this should be done in a vacuum desiccator over sulphuric acid, or at a temperature not exceeding 70°. For detection of caramel in milk, vinegar, and liquors, special tests are given elsewhere.

Fradiss Test.*—The dried residue of the sample to be tested is extracted with warm, pure methyl alcohol, which, if caramel be present, is colored brown. Filter, and to the filtrate add amyl alcohol or chloroform. In presence of caramel, a brown flocculent precipitate is formed, which slowly settles to the bottom of the tube.

Indigo in aqueous solution turns green with ammonia. On boiling, the solution becomes bright blue. Indigo in neutral or acid solution dves wool or silk.

ANIMAL COLORS.

Cochineal.—This dyestuff is used in ketchups, cordials, confections, and other food products. Robin's test for cochineal is as follows: The aqueous solution is acidulated with hydrochloric acid, and shaken out in a separatory funnel with amyl alcohol. Cochineal imparts to this solvent a yellowish color, the depth depending on the amount present. The separated amyl alcohol is washed with water till neutral, and divided into two portions. To one of these a little water is added, and then drop by drop a solution of uranium acetate, shaking each time a drop is added. In presence of cochineal the water is colored a very characteristic emerald-green color. To the other portion ammonia is added. If cochineal has been used, a violet coloration is produced.

MINERAL PIGMENTS.

Evidence of the presence of these pigments is usually best looked for in the ash of the suspected sample. In some cases the color may be extracted from the dried residue by water, alkali, or alcohol.

Prussian Blue.—This pigment is insoluble in water. It is decomposed and decolorized by treatment with potassium hydroxide. If the

^{* ()}estr. ungar. Zeits. Zucker. Ind., 1899, 28, 229-231; Abs. Zeits. f. Unters. Nahr. u. Genuss., 2, 1899, p. 881.

filtered alkaline solution of the coloring matter be treated with hydrochloric acid and ferric chloride, a precipitate of the original Prussian blue will be produced. For reactions on the fiber see table, page 810.

Ultramarine Blue is decolorized by hydrochloric acid with evolution of hydrogen sulphide, which blackens filter-paper moistened with lead acctate. For the recognition of ultramarine in sugar see page 613. For its detection on the fiber see table, page 811.

Chromate of Lead has never been used to any extent in food products with the exception of confectionary. For its detection, see page 647.

COAL-TAR COLORS.

So many of the coal-tar dyes are adapted for use in food that it would be impossible to even name them all, especially in view of the fact that new colors are from time to time being added to the list. No attempt will be made in the present work to give the nature and composition of the dyes named, as such descriptions would lead beyond its scope. For detailed information along this line the reader is directed to the references on page 813, and especially to the works of Schultz and Julius, Benedict and Knecht, Weyl, etc.

About 2000 separate coal-tar dyes are at present on the market. Various classifications of these colors are attempted, based on (1), their origin, as anilin dyes, naphthalin dyes, anthracene dyes, etc.; (2), their chemical composition, as nitro, nitroso, azo, diazo, and other compounds; (3), their solubility in water and other solvents; and (4), their mode of application to the fiber, as basic dyes, acid dyes, direct cotton dyes, mordant dyes, etc.

These dyes are sold in the form of powder, and are readily made into solutions for food colors in the case of the water-soluble varieties, and into pastes in the case of the insoluble forms. Most of the coal-tar colors employed in foods are naturally of the soluble variety, especially such as are found in jellies, jams, fruit products, canned foods, ketchups, beverages, and milk. Pastes made from insoluble dyes are adapted mainly for exterior coatings of hard substances such as candies. Colors in the dry form are to be looked for in such spices as cayenne, mustard, and mace, but a commoner method of coloring these spices high in oil is to mix with them a solution of the color in oil (usually cottonseed). Oil solutions of coal-tar dyes are also employed for coloring butter and oleomargarine.

The chief concern of the food analyst, as regards artificial color is

its recognition in food products. Coal-tar dyes may usually be identified as such, but it is not always possible to name the particular individual dye or combination of dyes employed, even though the class to which they belong may be determined. One reason for this is that not infrequently mixtures of two or more colors are employed.

Coal-tar Colors Allowed under the Federal Law.—The use of any dye, harmless or otherwise, to color food in a manner whereby damage or inferiority is concealed is in violation of Sec. 7 of the Food and Drugs Act of June 30, 1906. The addition of all mineral or metallic dyes, and of all coal-tar dyes, other than those specially provided for, is also prohibited. Pending further investigation the following coal-tar colors are permitted in foods, provided they are certified to be true to name and to be free from mineral and metallic poisons, harmful organic constituents, and contaminations due to improper or incomplete manufacture:*

Red Shades.—107. Amaranth [M.] [C.]. Synonyms: Fast red D. [B.]. Bordeaux S. [A.], azoacidrubine 2B. [D.], fast red EB. [B.].

56. Ponceau 3R. [A.] [B.] [M.]. Synonyms: Ponceau 4R. [A.], cumidin red, cumidin ponceau.

517. Erythrosin [B.][M.][B.S.S.]. Synonyms: Erythrosin D. [C.], erythrosin B. [A.], pyrosin B. [Mo.], iodeosin B., eosin bluish, eosin J. [B.].

Orange Shade.—85. Orange I. Synonyms: Alphanaphthol orange, naphthol orange [A.], tropæolin ooo No. 1, orange B. [L.].

Yellow Shade.—4. Naphthol yellow. S. [B.]. Synonyms: Naphthol yellow, acid yellow S., citronin A. [L.].

Green Shade.—435. Light green S. F. yellowish [B.]. Synonyms: Acid green [By.] [M.] [T.M.] [O.], acid green extra conc. [C.].

Blue Shade.—692. Indigo disulphoacid. Synonyms: Indigo carmine, indigo extract, indigotine [B.], sulphonated indigo.

None of these seven colors is patented, hence their manufacture is not likely to become a monopoly. They may be used in combinations, thus securing any desired shade. For example, violet may be obtained by mixing indigo disulphoacid and one of the red colors, a blue-green by mixing indigo-disulphoacid with naphthol yellow S. or light-green S.F. and so on.

^{*}The numbers preceding the dyes are those given in Green: A Systematic Survey of the Organic Colouring Matters founded on the German of Schultz and Julius, Londons 1904; the letters in brackets represent the manufacturers who originated the names.

DETECTION OF COAL-TAR COLORS IN FOODS.—There are various methods for the separation of coloring matters from food products, and these may be divided into three general classes: First, dying silk or wool with the color by boiling the fiber in a solution of the sample to be examined; second, extracting the color from a solution of the sample by the use of an immiscible solvent; and third, extracting the color from the dried residue of the sample by means of a suitable solvent. Of these the method of dying wool lends itself most readily to the analyst's use, by reason of its simplicity, and from the fact that almost without exception coal-tar dyes adaptable for food colors are substantive dyes, being readily taken up by wool.

Basic and Acid Dves.—The soluble coal-tar dyes are either basic or acid. Basic dyes are precipitated from their aqueous solution by tannin. Acid dyes are not so precipitated. Theoretically, all the basic colors are taken up by wool from a faintly alkaline or neutral bath, while the acid colors are left in solution. Thus if a dilute solution of the color be made faintly alkaline with ammonia and boiled with the wool, only basic colors will be taken up. If both acid and basic dyes are present in the same solution, the basic color should first be exhausted by the use of fresh pieces of wool in the ammoniacal solution, till they no longer take out color, after which the solution should be made slightly acid with hydrochloric acid and again boiled with wool, which under these conditions takes out any acid colors. Comparatively few basic colors are employed in foods. Basic colors can be removed from the fiber by boiling with 5% acetic acid. Acid colors are removed therefrom by boiling with 5\% ammonia. Having dissolved the dye from the fiber by the appropriate solvent as above, the decolorized fiber may be removed, and the solution evaporated to dryness on the water-bath. The residue consists chiefly of the dyestuff, and may be put through various reactions for identification according to Rota's scheme, page 707.

Methods of Dyeing Wool from Food Products.—The wool employed should be white worsted, or strips of white cloth, such as nun's veiling or albatross cloth. Care should be taken that the color is pure white and not the more common cream white. The woolen material should be freed from grease by boiling first in very dilute soda solution and finally in water. Strips of the woolen cloth, or pieces of the worsted thus previously cleansed, are boiled in diluted filtered solutions of jams, jellies, ketchup, fruit and vegetable products, and similar food preparations, or

in solutions of candy colors, or in wines, the clear solution of the sample to be tested being slightly acidified with hydrochloric acid.

Arata * recommends boiling the wool in a dilute solution of the food material to which potassium bisulphate has been added, using 10 cc. of a 10% solution of the bisulphate to 100 cc. of the solution to be tested. If the color solution is neutral, the wool should first be boiled in this before acidifying, to separate out any basic dyes. The dyed wool, after removal from the solution, is boiled first in water, and afterwards preferably in an alkali-free soap solution. It is then washed and dried. The dried fiber may then be subjected to the various reactions given in the table, pp. 804-811; for recognition of the dye, this method of identifying colors by means of reactions on the dyed fiber being one of the most convenient.

Some of the vegetable dyes (including lichen colors), also cochineal, dye wool directly, and these may be identified by reactions given in the table with the coal-tar dyes. Other vegetable colors, and the natural colors of fruits nearly always give a slight dull coloration or stain to the wool, but this is not, as a rule, to be mistaken for the vivid hues of the coaltar dyes. Moreover most of the vegetable colors on the fiber turn green when treated with ammonia. Care should be taken to thoroughly wash the wool after the dyeing, so that colored particles simply held thereon mechanically may be removed.

Sostegni and Carpentieri† recommend a method of double dyeing, applicable when acid dyes are employed. The method consists in first boiling the wool in a dilute acid solution of the food sample as above described, after which the fiber is removed and boiled, first in very dilute hydrochloric acid solution, and then in water, till free from acid. The color is then dissolved from the fiber by boiling the latter in a weak ammoniacal solution, some of the colors being more readily dissolved than others. The fiber is then removed from the solution, the latter is acidified with hydrochloric acid, and the color fixed on a fresh piece of wool by boiling therein. The second dyeing fixes coaltar and lichen colors on the fiber, but fruit colors and most others of vegetable origin remain in solution after this treatment. Any color left on the first fiber, after treatment with ammonia, is probably due to the

^{*} Ztsch. anal. Chem., 28 (1889), 639.

[†] Ibid., 35 (1896), 397.

natural vegetable color of the sample, and is usually no more than a dull stain.

Vegetable Colors on Wool.—In case no color is directly fixed on the fiber by boiling wool in a solution of the sample, either neutral or acid, absence of coal-tar colors may be assumed. In this case it is sometimes advisable to boil strips of previously mordanted white cotton in an acid solution of the sample, to remove certain vegetable colors for purposes of testing on the fiber. The cotton is mordanted by boiling in a dilute (5%) solution of potassium bichromate.

Extraction of Colors from their Solution by Immiscible Solvents.— Methods based on this principle are in use in the municipal laboratory at Paris.* Sanglé-Ferrière uses the following method: 50 cc. of the wine or solution to be tested for color are rendered slightly alkaline by ammonia, and cautiously shaken with about 15 cc. of amyl alcohol. If acid dves are present, they will be dissolved, and will impart to the amyl alcohol a distinct color.† Basic dyes also are dissolved, but when they are present the amyl alcohol solution is colorless. Remove the amyl alcohol by means of a separatory funnel, wash with water, and finally, if the alcohol is colored, dilute with about an equal volume of distilled water and evaporate on the water-bath with a piece of white wool. The wool should be kept in the solution till the odor of the amyl alcohol has disappeared, and, if not then colored, for a short time longer, as with some colors the wool will dve more readily in the aqueous solution than in the amyl alcohol. Remove the wool, and evaporate the solution to dryness. Test for color in the dried residue, and on the fiber also.

Orchil, like the acid colors, is extracted by, and imparts a coloration to the amyl alcohol under the above conditions, the color being a light violet.

If the amyl alcohol extract after separation, washing, and filtering is colorless, acidify with acetic acid; if a basic color is present, it will be indicated by a coloration at this stage; if there is no coloration on the addition of acetic acid, no basic color is present excepting fuchsin, which is separately tested for. In case a basic dye is indicated, add distilled water and evaporate with wool as before. Test the dried residue with pure concentrated sulphuric acid.

^{*} Girard et Dupré, Analyse des Matièrs Alimentaires, pp. 167, 581.

[†] Acid fuchsin forms an exception to this rule by dissolving colorless like basic dyes. A special test is, however, given for it, p. 797.

Fuchsin is indicated by a yellow-brown color with sulphuric acid, which by dilution with water becomes rose; sajranin, by a green color becoming first blue, then red, when diluted with water, and magdala red by a dark blue color, turning red on the addition of water.

Basic colors are also extracted readily, according to Robin, by making the solution to be tested alkaline with sodium hydroxide, and shaking with acetic ether. The solvent is removed, washed, and evaporated with wool (on which the tests are to be made), or the evaporation is carried to dryness and the tests made on the residue.

Many coal-tar colors are extracted by amyl alcohol in acid solution, but some of the natural fruit colors are also dissolved under these conditions. The coal-tar dyes thus dissolved will, however, dye wool and the fruit colors will not. Fruit colors are not extracted from acid or alkaline solution by ether, nor from alkaline solution by amyl alcohol.

Robin's method for ascertaining whether acid colors are present consists in adding to the liquid to be tested an excess of calcined magnesia, and a little 20% mercuric acetate solution, the mixture being boiled and filtered. If the filtrate is colored, or if by the addition of acetic acid to the colorless filtrate a color is developed, a coal-tar dye is indicated.

Separation of Acid and Basic Colors with Ether.*—Acid and basic colors may be separated from their dilute aqueous solution, according to Rota, by means of ether as follows: To 100 cc. of the solution add 1 cc. of 20% potassium hydroxide and shake in a separatory funnel with several portions of ether. Basic dyes are dissolved by the ether, leaving behind as a rule the acid colors.† Wash the ether extract with faintly alkaline water, and shake out with 5% acetic acid. Some colors remain in the ether, others are dissolved in the acid. Separate the two solvents, and evaporate each to dryness on the water-bath.

The acid colors left in the slightly alkaline, aqueous solution after removal of the basic colors by ether as above, may, if desired, be separated into several groups by successive extraction, as follows: first slightly acidulate with acetic acid and extract with ether, then acidify with hydrochloric acid and again extract, and finally examine the residual solution for colors that are insoluble in ether. Thus erythrosin and eosin are soluble in ether when shaken with their aqueous solution made acid with hydrochloric acid, while acid fuchsin is insoluble.

^{*} Analyst, 24, p. 45.

[†] A few acid dyes are exceptional in being soluble in ether with alkali, as for example, quinolin yellow and the sudans.

Separation of Colors from Dried Food Residues by Solvents.—This method is rarely employed, excepting in the case of colors insoluble in water, but soluble in ether or alcohol. The dried pulp of canned vegetables, ketchups, etc., may be acidified with hydrochloric acid, and the color extracted therefrom directly with alcohol. In this case however, there is no obvious advantage over the previous methods of dyeing the fiber directly in the acid solution of the sample.

Girard's Tests for Acid Fuchsin.*—Add 2 cc. of 5% potassium hydroxide to 10 cc. of the wine or other solution to be tested, or enough of the alkali to neutralize the acid. Then add 4 cc. of 10% acetate of mercury and filter. The filtrate should be alkaline and colorless. If the solution remains uncolored after acidifying with dilute sulphuric acid, no acid fuchsin is present. If, however, there is produced a red to violet coloration, and no other coal-tar colors have been found by the amyl alcohol extraction, the presence of acid fuchsin is shown.

Bellier's Test for Acid Fuchsin.—Presence of acid fuchsin is indicated by adding to 20 cc. of wine or other solution to be tested about 4 grams of freshly precipitated yellow oxide of mercury, boiling and filtering. The filtrate, if acid fuchsin is present, is colored red, tinged with violet.

According to Blarez, all red coal-tar colors, with the exception of acid fuchsin, and all red vegetable colors are completely decolorized by acidulating their aqueous solution with tartaric acid, and digesting with dioxide of lead.†

Schemes for Identification.—These serve for identifying unknown colors by their characteristic reactions, first grouping them into classes, and finally ascertaining the particular color itself. Of these may be mentioned the tabular schemes of Witt,† Weingartner,§ Green, Martinon, and Rota.**

Rota's Scheme is one of the latest, and on some accounts the best, being based on the relation between the color and the composition of

^{*} Analyse der Substances Alimentaires, p. 169.

[†] Allen, Commercial Org. Analysis, vol. III, p. 283.

[†] Zeits. anal. Chem., 1887, 26, p. 100; Analyst, 11, p. 111.

Jour. Soc. Dyers, etc., III, p. 67.

Jour. Soc. Chem. Ind., 12, No. 1.

[¶] Jour. Soc. Dyers, 3, p. 124.

⁺ Chem. Zeit., 1898, pp. 437-442; Anal., 24, p. 41.

the dyes. The colors are divided into two main groups, according to whether or not they are reducible by stannous chloride. These two groups are each further subdivided into two classes, the reducible colors being classed according to whether the color remains unchanged, or is restored by treatment with ferric chloride, and the non-reducible colors according to their action with potassium hydroxide.

The tests are carried out on a dilute aqueous or alcoholic solution of the coloring matter, the strength being about 1 in 10,000. Treat about 5 cc. of this solution with 4 or 5 drops of concentrated hydrochloric acid and about as much stannous chloride in a test-tube, shake the mixture, and heat if necessary to boiling. With some colors the process of decolorization is a slow one, especially if the solution is too concentrated, and it is well to repeat the experiment, if in doubt, diluting the original sample still further with water. Tin in solution in concentrated hydrochloric acid may be employed instead of stannous chloride, if desired.

Here, as in all cases of color testing, it is well to make comparative tests with known colors.

CLASSIFICATION OF ORGANIC COLORING MATTERS.

[A portion of the aqueous or alcoholic solution is treated with HCl and SnCl₂.]

Complete decolorization. Reducible coloring matters. Colorless solution is treated with Fe ₂ Cl ₆ , or shaken with exposure to air.		alone. Nonreducible	further than with HCle colors. A part of nixed with 20% KOH
The liquid remains unchanged. Coloring matters not reoxidizable.	The original color restored. Reoxidizable coloring matters.	Decolorization, or a precipitate. Imido- carbo-quinone color- ing matters.	
CLASS I.	CLASS II.	CLASS III.	CLASS IV.
Nitro, nitroso, and azo colors, including oxyazo and hydrazo colors. Picric acid, naphthol yellow, ponceau, Bordeaux, and Congo red.	Indogenide and imido- quinone coloring matters, methylene blue, safranin, in- digo carmine.	Amido-derivatives of di and triphenyl methane, auramins, acridins, quinolins, and color derivatives of thio benzenil. Fuchsin, rosanilin, auramin.	methane, oxy-ke- tone, and most of natural organic col-

CHARACTERISTICS OF ORGANIC COLORING MATTERS. CLASS I.—REDUCED BY HCl+SnCl, AND NOT REOXIDIZABLE.

Witro-coloring matters. R-NO.

Yellow or orange, soluble in ster. Wool and silk dyed directly, but not cotton. The aqueous solution shows tendency to decolorization with HCL With HCI+SnCL partially reduced. water.

Nitramines; soluble in ether in the presence of KOH.

-N-R-N(0. c.g., Aurantia.

Nitro - phenols; insoluble in ether

soluble in ether in O-R-N(O. presence of acetic acid, Nonsulphonated; in the presence of KOH,

Sulphonated; insoluble in ether.

Naphthol yellow.

Victoria yellow.

Nitroso-coloring matters, 0-R-N-OH Brown or green, usually insol-ole in water; indirect for fibers; ith H₂SO₄ + CHOH give blue with H₂SO₄+CHOH give bh color (Liebermann's reaction). uble in

A so-coloring matters.

R-N=N-R

Their aqueous solution decomposed with KOH and extracted with ether gives an etheral extract with annexed characteristics.

Nonsulphonated; insoluble in water; soluble in alcohol: soluble in ether in Dioxin (L). presence of acetic acid.

Nonsulphonated Sulphonated; soluble in water; insoluble in ether.

ing matters.
Oxyazo coloring
matter, without amido-azo color-Colored, shaken with dilute acetic acid yields to it the original color. Basic coloring matters.

Colored solution; not yielding its color to dilute acetic acid. Neutral coloring matters,

carboxyl.

Non sulphonat-

ed; extracted by ether from dilute not extracted by solution in acetic Sulphonated; ing to aretic acid. Acid coloning mattion; yields noth-Colorless

boxyl group.

Direct for cotton | Chrysamin. Indirect for cot- \ ton wool ton wool, ton wool WOOL. Oxyazo coloring matters, with carpounds; unaltered Nonamido com-

Naphthol green.

-N-R-N-NHR, Bismarck brown.

Sudan 1 (A).

Indirect for cot- Diamond yellow (By), O-R-N-HRy.

Solid yellow N (P). Indirect for cot- | Bordeaux B (A). Direct for cotton { Congo red (A). Direct for cotton Azo blue (A).

by HNO.

ether from solu-

tion in dilute acetic

younds; changed Amidocom by HNO.

wool.

REONIDIZABLE.
AND
+SnCl,
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REDUCED
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LASS

E. Types.	N Nile blue A (B).	$\begin{pmatrix} R_1 \\ N \\ R_{-N} \end{pmatrix}$ Methylene blue.	$\begin{pmatrix} R_{1} \\ N \\ N \\ R = N = \end{pmatrix}$ Indulin soluble in alcohol.	$N \longrightarrow N = \begin{cases} N \longrightarrow N = \\ N & \end{cases}$ Safranin T extra (A).	$\begin{bmatrix} R \\ R = O \end{bmatrix} \text{Indophenol.}$	NH CO Indigotia.	N O Fluorescent blue; orcein.	Sulphonated indogenides. Indigo carmine. Sulphonated thiazins. Thiocarmine R (C)
CLASS IIKEDUCED BY HCI+ShC1, AND KEONIDIZABLE.	Oxyazines (no sulphur).	Thiazins (sulphur).	Indulins; blue color with concentrated H _z SO ₄ . Blue dilution.	Safranins; green color with H ₂ SO ₄ . On dilution blue, then violet.	Indophenols.	· Indogenides.	Oxazonea.	- ~- ;
CLASS II.—KEDUCED BY	solution is readily by HCl+SnCl, in	the cold,	The colored solution is reduced but slowly and in-	completery, even on warming, and with the addition of much SnCl ₂ +HCl.	Blue coloring matters changed by HCl on warm- Indophenols, ing.	Red or blue coloring mat- ters. Unaltered by HCl. With HNO ₂ yield isatin.	Nonsulphonated. Solu- ble in ether in presence of acetic acid.	Sulphonated. Insoluble Reduced by SnCl,+HCl. in ether under all circum-
		The ethereal solution is colored or colorless, and yields the original color to	5% acetic acid. Basic coloring matters fixed on wool in alkaline bath.		Colored; does not yield the color to acetic acid. Neutral coloring matters.	Insoluble in water. Soluble in alcohol. Fixed on fibers in bath.	Uncolored; yields noth- ing to acetic acid.	_

The aqueous or alcoholic solution is treated with KOH and extracted with ether.

The ethereal solution washed with water has the annexed characteristics.

CLASS III.—COLORING MATTERS NOT REDUCED BY SnC2+HCl. CONTAINING THE IMIDO-QUINONE CARBON CHROMOPHORE -N-R-C-, CHARACTERISTICS OF ORGANIC COLORING MATTERS—(Continued).

HORE -N-R-C	$\left. egin{aligned} Tytes. \ ext{E.g. Auramin O (B).} \end{aligned} ight.$	Phosphin.	Fuchsin.	$\left. ight\}$ Pyronin (G).	Rhodamin S (By).	Quinolin yellow A (soluble in alcohol).	quin- { Quinolin yellow A { (soluble in water).	$race{ \text{Fuchsin S (B).} }{ ext{Violamin R (M).} }$	Primulin (B).
NONE CARBON CHROMOI	C.R.R	a d	(nonsul- $\begin{cases} C & R \\ C & R = N - \end{cases}$	lored (—C O)	(non- Unal-	nes. $\begin{cases} -C \\ R = N = \end{cases}$	Sulphonated quin- one-phthalones, Sulphonated fuch-	sins. Sulphonated rhoda- Violamin R (M).	0-0 = -4
COLLEGIA MATTERS NOT KEDUCED BY SHC2+HCI. CONTAINING THE IMIDO-QUINONE CARBON CHROMOPHOREN-R-C	Auramins.	Acridins	Fuchsins (no phonated).	Pyronins (colored yellow by HCl. Direct for cotton wool).	Rhodamins (sulphonated. Itered by HCl).	Quinone-phthalones.	and alkalies. }	hanged by HCl. \(\) soluble in water \(\) HCl. Changed \(\)	f. Thiazoles.
	Colorless, nonfluorescent ethereal solution. Yellow color yielded to acetic acid nonfluorescent. The aqueous solution is decolorized by KOH and decomposed by HCI.	Colorless, ethereal solution. Green fluorescence. Aqueous solution precipitated by KOH, hardly altered by HCl. Turns red with HNOs.	Colorless, or colored ethereal solution. Non-fluorescent. Color yielded to acetic acid—freddish violet, blue, and green without fluorescence. Aqueous solution usually decolorized on warming with KOH, and colored yellow by HCl (excepting fuchsin).	Ethereal solution colorless and nonfluores- nt. Acetic acid colored rose and fluoresces.	Aqueous solution decolorized with KOH.	The ethereal solution is yellow, and nonfluorescent. Alcoholic solution yellow, nonfluorescent, and unaltered by aqueous acids and alkalies.	Yellow coloring matters. No fluorescence in water. Unaltered by aqueous acids and alkalies. Reddish violet, blue, or green coloring matters.	Usually decolorized by KOH, little changed by HCI. Red or violet coloring matters. Soluble in water with fluorescence. Precipitated by HCI. Changed. but little, or not at all, by KOH.	Brownish yellow or orange coloring matters. Aqueous solution ± fluorescent. Fixed directly on silk, wool, and cotton.
	Colorless Yellow colo cent. The	Colorless cence. Aqu hardly alter	Coloriess fluorescent. reddish vio rescence. ized on wal	Ethereal cent. Acel		solution o acetic s. In- in alco-	ot dye	Does no	Dyes the wool.
CHAIL COLORING MAIN		The color is yielded to 5% acetic acid. Basic coloring matters. Fixed on wool in alkaline bath (NH _s).			The colored ethereal solution does not yield its color to acetic acid. Neutral coloring matters. Insoluble in water. Soluble in alcohol.	Ethereal solution colorless. Yields nothing to acetic acid. A cid coloring matters. Soluble in water. Fixed on wool in acid bath (HCl).			

-R-C-	Types.	•	Alizarin yellow A (B).	etin.	.	Sulphonated alizarin (alizarin red).
ORE O	Aurin.	Eosin.	Aliza	Quercetin.	Alizarin	Sulph (ali
RBON CHROMOPHO	$\left\{\begin{array}{c} R_1 \\ C-R_1 \\ R=0 \end{array}\right\}$	$\left\{\begin{array}{c} C_{-R_1} \\ C_{-R_2} \\ \end{array}\right\}$	# 00 m			
HE OXY-QUINONE CA	Aurins.	Phthaleins.	Benzophenones.	Flavones.	Nonsulphonated anthraquinones.	Sulphonated an- thraquinones.
CLASS IV.—COLORING MATTERS NOT REDUCED BY SAC1,+ HCI. CONTAINING THE OXY-QUINONE CARBON CHROMOPHORE O-R-C-,	Not directly fixed on wool. Most of them insoluble in water. Soluble in alcohol without fluorescence.	Fixed directly on wool. ost of them soluble in ster and alcohol. Fuores-nce.	Inclined to decol- orization, especially on warming (with decomposition).	Colored intense yellow without decomposition.	The free coloring matter precipitated. Usually soluble in ether, and indirect for fibers.	Coloring matters remain in solution. Insoluble in ether, fixed directly on wool.
BY SaCi,	Not o Most o water.	Fixed Most c water a	kaline so- ated with	The alT ort noitul to seeses	kaline solu- dified with L	Is aTT iva noti iva nisse iva sitessit
ERS NOT REDUCED	The coloring mat-	suspended in boiling water.	Dissolves with	dishyellow color. Monoketones.	Dissolves with red, reddish violet, green, or hine color	Diketones (quinones).
ING MATT	<u>=</u>		ei 19 7 9 16W		autei uitm de	Theate
CLASS IV.—COLOR	Remains unaltered. Nonamido triphenylmethane coloring	matters. Usually uble in water and rect for wool.		Changes to green or olive green.	coloringmatters. Most of them insoluble in water and indirect for fibers.	
	MOI W II	(1:1000).	on of Fe _s Cl	on or the cok dilute solutio	s lo sqorb	r our

The alcoholic solution of the coloring matter treated with a few

Direct Identification of Colors.—In identifying the colors commonly used in food, it is rarely necessary to carry out such involved processes of analyses as are rendered necessary by Rota's scheme. It is frequently possible to ascertain the color or group of colors present by making direct tests with various reagents, either on the dyed fiber as described on p. 812, or on the dry coloring matter, or in a solution containing it.

Many tables for this purpose are prepared, but they are never complete by reason of the many new dyestuffs constantly introduced. Such tables are to be found in Allen's "Commercial Organic Analysis," and in Schultz and Julius's "Systematic Survey of the Coal Tar Colors." While it is true that the limitation of the dyes suitable for purposes of food coloring imposes a somewhat lighter task on the food analyst than that of the chemist who has to deal with all varieties of commercial colors, yet it is obviously impossible to make a complete list covering even the restricted field of food colors. Doubtless there are colors long well known that would serve admirably for this purpose, but have never yet been tried.

Mainly from such sources as the above comprehensive tables of colors and their reactions, the writer has compiled the table on pp. 804-811. taking as a basis the scheme of Allen.* This table includes over fifty selected coloring matters, which are adapted for, and have been found in, foods by various analysts, as listed in state and government reports, as well as in laws of various countries dealing with food colors. This table will at least contain the colors most commonly met with, and will nearly always serve, if not to identify the exact dye, to aid in classifying it. In case the analyst wishes to identify the color, he should be provided, for standards, with as complete a collection of known purity dyestuffs as possible covering the colors he is likely to meet with in foods, and should make comparative tests, if the slightest doubt exists. If the unknown color is apparently not found in the following table, and the more exhaustive tables are unavailable, it is still possible to locate the dye, by making similar tests on other standard colors suggested by the behavior of the unknown color, and carefully comparing them.

Most difficulty is encountered when the coloring matters are mixtures instead of simple dyes. In this case it may be necessary to resort to fractional extraction by ether, as suggested by Rota (p. 796), in order to separate the colors.

^{*} Commercial Organic Analysis, Vol. III, pt. 1, 3d ed., p. 530 et seq.

IDENTIFICATION OF COLORS ON THE FIBER.

<u>-</u>			cutaincie of Dyestuit.	Acid	Hydrochik	Hydrochloric Acid.	unudrac	Sulphuric Acid.
_		Dry.	Solution.	Basic.	Fiber.	Solution.	Fiber.	Solution.
RED COLORS.		Beetle-green	Red	Basic	Yellow*	Yellow	Brown * yellow	Yellow
	í	Green glisten-	Crimson	Acid	Paler	Pink	Paler	Pink
	(B.S.S.)	Brown powder	Red	Basic	Blue	Yellow	Green	Green
Eosin A (Carmoisin.	(B.)	crystals Red-brown	Crimson	Acid	renow Dark-red vio-	Lilac	venow Violet-black	renow Violet-black
Phloxin.	:	powder Brown-yellow	Red, greenish		let * Yellow		Yellow	
Erythrosin.	:	powder Brown powder	fluorescence Red		Orange		Yellow	Orange
Crocein scarlet 7B (F. Ponceau 6RB	(By.)	Reddish brown	Red	Acid	Violet †	Colorless ‡	Blue	Blue
	(K.)	Brown powder	Orange	Acid	Violet	Colorless ‡	Green	Green
-	(B)	Brown powder	Orange	Acid	Violet Black	Colorless ‡	Green	Green
	By.)	Red powder	Red	Acid	Dark-red violet	Lilac	Dark-red violet	Lilac
:::::::::::::::::::::::::::::::::::::::	B.)	Brown powder	Red	Acid	Decolorized		Brown	į
Croceine scarlet 3B (F. Ponceau 4RB	Ber.)	Brown powder	Ked	Acid	Blue	Blue	Bine	Blue
eau 3R.	£	Red powder	Red	Acid	Cherry red	Cherry red	Cherry red	Cherry red
Cochineal red A	(œ	Scarlet-red Dowder		Acid	Darker		Crimson	
:	B)	i	Reddish brown	Acid	Purple		Violet	Violet
Fast red B		Brown powder	Purple	Acid	Crimson	Pink	Violet	Purple
	(E) (E) (E) (E) (E) (E) (E) (E) (E) (E)	Brown powder Reddish-brown	Magenta-red Magenta	Acid	Darker Darker	Pink Pink	Violet Violet	Violet Violet
-		powder						

				A	IRTIFI	CIAL .	FOOD (OLO	ORS.					805
Alcebel	Aconor	Bluish red	Pink	Red fluoresc. Sol. cosin not extracted; spirit	No action Pink with green	No action	No action No action No action	Nc action	Pink	No action	Pink	Pink	Pink Pink	l d
Stannous	Chloride.	Decolorized *	Little change	Decolorized Yellow	Decolorized Yellow	Orange Brown	Decolorized Decolorized Black, then	Decolorized on	neaung Decolorized Decolorized	Decolorized on	Lighter	Lighter	Lighter Lighter	On standing liquid becomes blue or greenish blue. Spyes both cotton and wool in alkaline or neutral bath.
Spot with	Nitric Acid.	Yellow	Yellow	Blue Yellow	Yellow	Yellow-red rim Dark blue	Blue to brown Blue to brown Black	Brown	Yellow Yellow, blue	Yellow	Yellow Yellow	Yellow	Yellow Yellow	t On standing liquid becomes blue or greenish blue. S Dyes both cotton and wool in alkaline or neutral
Ammonia.	Solution.			Pink Yellow fluoresc.	Pink Pink	Pink	No action No action	Orange	Pink Pink	Pink	Pink		Pink Brown	On standing liquidates both cottor
Amn	Fiber.	Decolorized	Decolorized	Yellow	Unchanged Pink	Orange Blue	No action No action No change	Orange	No change	No change	Darker		Brighter Darker	
Sodium Hydroxide.	Solution.			Pink Pink fluoresc.	Pink Pink	Pink	Colorless Colorless			Brown	Brown			on washing.
Sodium F	Fiber.	Paler	Decolorized	Yellow	Brownish Pink	Pink Blue	Violet Violet No change	Orange	Darker Purple	Brown	Brown Maroon	Red	Brown	Color restored wholly or in part on washing.
		RED COLORS.	Acid fuchsin	Safranin. Eosin A.	Carmoisin.	Erythrosin	Fonceau or D. Biebrich scarlet Fast ponceau B.	Azo eosin.	Rose bengal	Cumidine red, Ponceau 3R	Cochineal red A. Fast red A.	Fast red B	Fast red C	* Color restored † On standing fi

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	11	IDENTIFICATION	N OF COLORS	RS ON	THE	FIBER—Continued.		
		Character or Dyestuff.	Dyestuff.	Acid	Hydrochloric Acid.	nic Acid.	Sulphuric Acid.	c Acid.
		Dry.	Solution.	Basic.	Fiber.	Solution.	Piber.	Solution.
RED COLORS—(Continued). Orchil substitute V.	(F.)	Brown paste	Red-brown	Acid	Crimson	Crimson	Crimson	Crimson
Scarlet 6R. Orchi *	(K)	Brown powder Dark-red paste	Purple Red	Acid	Crimson No change	Pink Red	Violet Purple	Violet Purple
Barwood.		Dark-red wood Dark-red wood Fiery-red pow-	Red Red Yellow-red		Dark red Redder Decolorized	Pink	Brown Red-brown Decolorized	Yellow Dirty brown
Cochineal §		der Dry pinkish	Deep red		Orange	Orange	Pink	Pink
Cudbear * YELLOW AND ORANGE	*	:		:	Lighter	Red	Dirty brown	
Colors. Pictic acid. Victoria yellow	4 9 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6	Yellow crystals Orange powder	Yellow Orange	Acid	Decolorized Decolorized	Yellow	Decolorized	
		Orange-yellow	Yellow	Acid	Decolorized	Colorless	Decolorized	
. vi		Ora	Yellow	Acid	Decolorized	Colorless	Decolorized	
J	(B.S.S.)	Red-brown	Orange-brown	Basic	Red	***************************************		Yellow
Fast yellow.	Ê	Yellow powder	Yellow	Acid	Red		Вгочп	
Orange I		Red-brown	Orange	Acid	Magonta	Magenta	Magenta	Magenta
Orange II.	(£)	Yellow-red	Orange	Acid	Violet	Violet	Violet	Violet
Orange III. Methyl orange. Tropsolin D.	(By.)	Orange-yellow powder	Orange	Acid	Red	Pink	Violet	Violet

	Sodium F	Sodlum Hydroxide.	Amm	Ammonia.			
	Piber.	Solution.	Piber.	Solution.	Spot with Nitric Acid.	Stannous Chloride.	Alcohol.
RED COLORS—(Continued) Orchil substitute V.	Maroon		Maroon		Yellow	Lighter	
	Brown Purple		Purple	Pink	Yellow	Decolorized	Purple
Archil orseille. Brazil wood †	Maroon Purple	Violet Colorless	Violet Purple	Violet Colorless	Yellow Olive	Violet No change	Pale yellow Red
Cochineal \$	rale yellow Violet	Purple Violet	Fink Violet	Purple Violet	Yellow Yellow, red	Yellow Orange Decolorized	No action No action Bluish red
YELLOW AND ORANGE COLORS, Picric acid.	Orange	Yellow	Paler	Yellow	臣	Decolorized	Yellow
Victoria yellow Anilin orange. Dinitrocresol.		4 a 4 5 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6	No change				
	Orange	Yellow	Paler	Yellow		Decolorized	Yellow
· s	Paler	Yellow	No change	Yellow	•	Bleached	No action
Chrysoidine yellow.	Paler No action		Yellower No action	# E	Red	Decolorized Decolorized	Yellow Pale yellow
Acid yellow. Orange I.	Deep red	Deep red	Deep red	Deep red	Orange	Decolorized	No action
Orange II.	Deep red		No action	•		Decolorized	No action
Orange III. Methyl orange. Tropseolin D.	Yellowish	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	No change	Yellow	4 4 4 9 0 0 1 4	Decolorized	Pale yellow
	a wool in neutral of a not dye wool unl	or slightly acid solves mordanted with	Does not dye wool unless mordanted with alum or chrome. Does not dye wool unless mordanted with alum or chrome. Does not dye wool but does ontoto in alleging columns.	•	Dyes wool by tin mordant. Warm water extracts color.	mordant.	

4 Does not dye wool, but dyes cotton in alkaline solution.

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	Sulphuric Acid.	Solution.	Viole	Dark red	Red	Blue	Brown		Yellow	Brown	Вгоwп	Yellow		Red	Violet	Purple Red	
	Sulphur	Fiber.	Violet	Brown	Orange-yellow	Blue	Вгоwп	Duller	Yellow	Brown	Вгомп	Bright yellow Brown	Dark purple	Crimson	Violet	Purple Reddish	No action
FIBER—Continues	Hydrochloric Acid.	Solution.	Violet	Brown	Red	Red	Colorless	Yellow	Yellow	Orange	Yellow	Yellow Yellow	Purple	Pink		Yellow Red	
THE	Hydroch	Fiber.	Violet	Brown	Red	No change	Brown	Decolorized	No change	Orange	No change	No change No change	Purple	Scarlet	Violet	Darker Reddish	Decolorized
RS OF	Acid	Basic.	Acid	Acid	Acid	:	:	:	:	:	i		Acid	Acid	Acid	Acid Basic	
IDENTIFICATION OF COLORS ON	Character of Dyestuff.	Solution.		Pale yellow	Orange-yellow	Cream	Yellow	Yellow	Yellow		Yellow	Pale yellow Yellow	Orange	Reddish yellow	Yellow	Brown Brown	Brown
ENTIFICATION	Character	Dry.		Dull-yellow	powder Fiery-red	Brown-red	Dark-yellow	Yellow heavy	Yellow or buff	powder Yellow Brazil	Small bits of	Yellow-green	ury bernes Brownish-yel- low powder	Yellowish-red	Brown powder	Brown powder Brown powdér	
Ħ	Ä			(C.) (B.S.S.)	(By.)								(0)	(H.)		3	
			YELLOW AND ORANGE COLORS—(Continued). Orange IV.	Tropæolin OO. Primulin orange.	Crocein orange.	Annatto *	Turmeric	Chrome yellow †	Quercitron bark ‡	Old fustic ‡	Young fustic ‡	Weld ‡Persian berries ‡	Metanil yellow. Tropæolin Victoria vellow	Orange G.	Chrysamine R	Acid brown G. Bismarck brown.	brown §

	Sodium	Sodium Hydmxide.	Y W	Ammonia.			
					Spot with	Stannous	Alcohol.
	Fiber.	Solution.	Fiber.	Solution.			
YELLOW AND ORANGE COLORS							
Orange IV.	No action		No change	Yellow		Decolorized	Yellow
Primulin orange.	Dark brown		No action		Brown	Decolorized	Pale orange
Crocein orange.	Browner		Slightly darker			Decolorized	
Annatto *	No action		No action			Decolorized	Yellow
Turmeric.	Brown	Brown	Brown	Orange	Yellow	Brown	Yellow
Chrome yellow 7	Kedder	Yellow	No action			Decolorized	No action
Quercitron bark ‡	No change	Yellow	No change	Yellow	Blue	No change	No action
Voing fisher +	No change Brown		No change	xellow.	Pale yellow	No action	No action
Weld	No change	Vellow	No action		No action	No change	No action
Persian berries ‡	No change	Yellow		Pale yellow	Brown	Brown	No action
Metanil yellow.	Brighter		No change		Red	Brown	Pale yellow
Tropæolin.							
Orange G.	Terra cotta		No change		Yellow	Decolorized	
Chrysamine R	Darker		Darker	Pink	Red		Yellow
BROWN COLORS.							
Acid brown G	Darker		Brighter	Brown	Yellow	No change	Brown
Vesuvine	renower	Coloriess	No change	Brown		Decolorized	rink
Manganese brown §	No action		No action			Decolorized	No action

Kare(CN),—blue. Has—black

Fe₂Cl₆—olive. # Ash contains Mn.

DENTIFICATION OF COLORS ON THE FIBER—Continued

	j	Character of Dyestuff.	f Dyestuff.	Acid	Hydrochl	Hydrochloric Acid.	Sulphuric Acid.	c Acid.
		Dry.	Solution.	Basic.	Fiber.	Solution.	Piber.	Solution.
VIOLET-BLUE AND GREEN								
Methyl violet		Green metallic	Violet	Basic	Вгожп	Amber	Brown	Amber
Azo blue. Indulines soluble.	'Ву.)	Black powder Bronze powder	Violet Violet		Black Violet	Colorless Blue	Black	Blue Dark blue
Indigo. Indigo disulphoacid		Indigo powder	Indigo Indigo	Acid	No change Indigo	Bluish	Indigo	Blue Bluish
Indigo carmine. Prussian blue *	:	Deep-blue powder	Blue	:	No action		No action	Blue
Ultramarine blue † Methylene blue ‡ Resorcin green *	(B.) (D.H.)	Bronze powder Gray-brown	Blue Green		Decolorized Decolorized Yellow	Blue-green	Decolorized Green	Green
Malachite green New Victoria green.		Green crystals	Bluish green	Basic	Orange	Orange	Bleached	Orange
Naphthol green	(C)	Dark-green	Green		Yellow	Yellow	Olive	Drah
Light green SF, yellowish	(B.)	Dark-green powder	Green		Orange .	Yellow	Orange	Yellow

	Sodium H	Sodium Hydroxide.	Апп	Ammonia.		i	
*	Fiber.	Solution.	Piber.	Solution.	Nitric Acid.	Chloride.	Alcohol.
VIOLET-BLUE AND GREEN							
Methyl violet	Decolorized	Coloriess	Lilac			Green	Violet
Azo blue. Red Indulines soluble.	Red	Pink Violet decolor-	Dark violet	Red AsNaOH	Dark green	Decolorized Violet solution	No action Blue-violet
Fast blue R.	1	ized by zinc dust					
Indigo. Indigo disulphoacid	No change Olive	Yellow	No change Green	Yellow	Yellow	Paler, solution Decolorized	Blue No action
Indigo carmine.	P		1				W
Ultramarine blue †	No action		No action		uaero	Decolorized	No action
Methylene blue ‡	Violet		No action		Green	Decolorized	Greenish blue
Resorcin green *	Darker	Greenish	No action		Brown	Paler	No action
Malachite green	Decolorized		Decolorized		4	Decolorized	Green
New green.	Brichter	o di	No ention		200	Decolorised	Data hine
Light green SF, yellowish	Decolorized		Decolorized	4	Yellow	Lighter	No action

ABBREVIATIONS OF PRIME.

CaOCIg-decolorised.

Ash contains Pe.

ller, London. Paris.

Reagents.—In applying tests on the fiber, the reagents commonly used are as follows: Concentrated hydrochloric acid, concentrated sulphuric acid, sodium hydroxide (10% solution), strong ammonia (28%), a hydrochloric acid solution of stannous chloride, and concentrated nitric acid. The tests should be made on pieces of the fiber in small porcelain evaporating-dishes, which more readily than test-tubes show exact shades of color. In cases of suspected fluorescence, test-tubes should be used. Nitric acid is conveniently applied by a glass rod to the fiber. The stannous chloride should first be allowed to act in the cold. If no change occurs, gentle heat should then be applied, and finally boiling.

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CHAPTER XVIII.

FOOD PRESERVATIVES.

Preservation of Food.—Various processes have from ancient times been known and used for arresting the fermentative changes which food products in their natural state undergo on long standing. These processes include pickling with vinegar, drying, smoking, salting, preserving with sugar, and finally in the employment of heat in sterilizing and pasteurizing, and of low temperature as in cold storage. All of them are still in use, and are universally regarded as unobjectionable. In addition to these old and well-known methods of food preservation is the comparatively modern practice of arresting fermentation by the use of such antiseptic chemical agents as formaldehyde, beta-naphthol, boric, salicylic, benzoic, and sulphurous acids or salts of these acids, etc., in regard to the wholesomeness of which there is considerable difference of opinion. These substances depend for their efficiency on the more or less complete inhibition of bacterial growth. Nearly all exert a powerful antiseptic influence, to such an extent that to accomplish their object only small quantities need be used in food.

Apart from their toxic effects, a marked difference naturally exists between the employment of such substances as salt, sugar, and vinegar for food preservation, all of which are in themselves foods, and in the use of chemical agents that have no food value. The advocates of the use of chemical antiseptics claim that there are no authentic instances on record of injury from the use of such small quantities of these substances as are necessary to arrest decay, while there are many cases of injury arising from the use of foods which, while apparently wholesome, have undergone such fermentation as to develop ptomaines or other harmful toxins, and that because antiseptics prevent such spoiling of the food, their use is decidedly beneficial; that there is, besides, no more reason why a prejudice should exist against the employment of these

newer chemicals than against saltpeter, which has long been used in the corning of meat, or against the cresols and phenols left as a product of smoking.

The opponents to their use assert, that the addition to food of such antiseptic substances as prevent its decay also serves to retard the digestive processes when the food is eaten; that many of these substances are drugs, and as such cannot fail even in small quantities to exercise a toxic effect of some sort on the system; that finally their use is objectionable, as allowing the employment in certain foods of old materials that have in some cases already undergone incipient decomposition before the addition of the antiseptic, and are thus unwholesome.

Regulation of Antiseptics in Food.—In the absence of legislation directly prohibiting the use of any of the above-named antiseptics, and in view of the difference of opinion regarding their toxic effects when present in small quantities, it is difficult to maintain a complaint under the general food laws as they exist in most states, basing the complaint solely on their harmfulness. In some localities certain antiseptics are specifically allowed and others are prohibited. Some of the states, as, for example, Massachusetts, have special laws under which it is required that in the case of all foods thus treated, the name and percentage of such antiseptics as are used must appear plainly on labels of the packages or containers thereof, such a provision being based on the assumption that the general public should be informed of what they are buying, where any doubt exists as to the wholesomeness of any ingredient present. Where such laws as these are in force, the chemist's task is comparatively easy, in that conviction in court is not dependent on his individual opinion regarding the toxic effects of the antiseptic employed.

Physiological experiments for testing the toxicity of these chemical preservatives were formerly confined to the lower animals, but no satisfactory results could be thus obtained. Later, metabolism experiments were made on human beings treated with varying amounts of the preservatives under carefully controlled conditions, but the results of these, though made by experts of unquestioned ability, do not agree. Even if any of these substances as used in food appear to have little or no effect on people in good health, they cannot be assumed to be equally harmless to those who are inclined to be delicate or sickly. Even though pronounced harmless in themselves, there is still the objection that the chemical preservatives may readily conceal unclean methods or materials. If perishable foods are free from preservatives and are sweet and

untainted, the consumer has reason to believe that clean and wholesome materials and sanitary processes were employed throughout in their manufacture.

Commercial Food Preservatives.—A large number of commercial preparations are sold for purposes of preserving specific articles of food and are put out under trade names that usually convey no suggestion of their true character. Some of these consist of a single antiseptic substance, such as salicylic acid, ammonium fluoride, calcium sulphate, borax, or benzoic acid, while others are mixtures of several antiseptics, of which the following are typical examples, showing their composition as found, together with the amount of the mixture to be employed.

A. For preserving sausage meat, using 8 ounces per 100 pounds of meat:

Borax	36%
Salt	46%
Saltpeter	18%
(Colored with an anilin dye.)	

- B. For preserving cider and ketchup.
- A 34% solution of beta-naphthol in alcohol, using 2 fluid ounces to 45 gallons of cider, or 1½ ounces to 10 gallons of ketchup.
 - C. For preserving beer, using 11 ounces per barrel of beer:

Salt	45%
Salicylic acid	27%
Sodium carbonate and salicylate	28%

D. For preserving chapped meats, using 1 ounce to 50 pounds of meat.

Sodium sulphite	65%
Borax	35%

E. Effec.ive for curing beef, hams, tongues, bacon, pig's feet, etc.:

Borax	28%
Boric acid	12%
Sodium chloride	35%
Potassium nitrate	25%

F. For preserving milk and cream:

Boric acid	75%
Borax	25%

G. For preserving jellies, jams, preserves, mince-meat, and syrups, using from 1 to 2 ounces of preservative to 100 pounds of product:

Sodium benzoate	50%
Boric acid	40%
Sodium chloride	5%
Sodium bicarbonate.	5%

H. For preserving ketchup and tomato pulp, using from 6 to 8 ounces to 45 gallons of the product:

Sodium benzoate	50%
Sodium chloride	40%
Sodium sulphite	10%

I. Effective for keeping butter from becoming tainted or rancid, also for salt codfish, using 8 to 12 ounces per 100 pounds butter:

Boric acid	25%
Borax	50%
Sodium chloride	25%

J. For preserving eggs (surface application). A saturated solution of salicylic acid in 3 quarts of water, 1 quart strong alcohol and 7 ounces of glycerin.

FORMALDEHYDE.

Formaldehyde (HCHO) is a gas formed by the action of a red-hot spiral of platinum wire on vaporized methyl alcohol. It is also produced by the dry distillation of calcium formate. In the market it commonly appears in the form of a 40% solution of the gas in water under the name of formalin, and for use as a food preservative dilute solutions of from 2 to 5 per cent strength are usually employed. Its use as a food preservative is comparatively modern. Formaldehyde, while not confined exclusively to milk products, is, as a matter of fact, more commonly used in these than in other foods. Its prompt and direct action in checking or preventing the growth of lactic acid bacteria renders it especially desirable for use as a milk and cream preservative, from the standpoint of the dairy man who does not concern himself as to whether or not its use is injurious or illegal.

When present in milk to the extent of 1 part formaldehyde to 20,000 parts milk (a proportion quite commonly employed), the sample is kept

sweet for four days in summer weather, when under ordinary conditions, the milk untreated would curdle in less than forty-eight hours.

Determination of Formaldehyde in the Commercial Preservative.—
(1) Iodometric Method.*—Mix 10 cc. of the aldehyde solution (diluted if necessary to a strength not exceeding 3% of formaldehyde) with 25 cc. of tenth-normal iodine solution, and add drop by drop a solution of sodium hydroxide, till the color of the liquid becomes clear yellow. The solution is set aside for at least ten minutes, after which hydrochloric acid is added to set free the uncombined iodine, and the latter is titrated back with tenth-normal thiosulphate. Two atoms of iodine are equivalent to one molecule of formaldehyde, in accordance with the following reactions:

$$6NaOH + 6I$$
 = $NaIO_8 + 5NaI + 3H_2O$.
 $3CH_2O + NaIO_8$ = $3CH_2O_2 + NaI$.
 $5NaI + NaIO_3 + 6HCl = 6NaCl + I_6 + 3H_2O$.

(2) Method of Blank and Finkenbeiner.†—Three grams of the solution are weighed into a tall Erlenmeyer flask, to which is then added from 25 to 30 cc. of twice-normal sodium hydroxide. 50 cc. of pure 2.5 to 3 per cent hydrogen peroxide solution are next gradually run in during a space of from three to ten minutes, through a funnel placed in the neck of the flask to prevent spurting, and the solution is allowed to stand for two or three minutes, after which the funnel is washed with water.

Finally the unused sodium hydroxide is titrated with twice-normal sulphuric acid, using litmus as an indicator. The less formaldehyde in the sample, the longer the mixture should stand after addition of the hydrogen peroxide, to complete the reaction. When less than 30% is present, it should stand at least ten minutes.

Ascertain the percentage of formaldehyde, by multiplying by 2 the number of cubic centimeters of soda solution used, when 3 grams of the sample are taken.

(3) Ammonia Method.‡—Weigh 10 grams of the formaldehyde solution into a flask, and treat with an excess of ammonia. Cork the flask and shake frequently during several days. The formaldehyde is by this process converted into hexamethylamine.

Transfer the solution to a tared platinum dish, and evaporate nearly

^{*} Zeits. anal. Chem., 1897, 36, pp. 18-24; abs. Analyst, 22, p. 221.

[†] Ber., 31 (17), 2979.

Conn. Exp. Sta., Annual Report, 1899, p. 143.

to dryness on the top of a closed water-bath. Finally the dish is transferred to a desiccator, and the drying continued over sulphuric acid to constant weight. The per cent of formaldehyde is calculated from the weight of the hexamethylamine, making a correction for the residue left by the formaldehyde itself by direct evaporation:

$$6CH_2O + 4NH_4OH = (CH_2)_6N_4 + 10H_2O$$
.

Or an excess of a standardized ammonia solution may be added in the first place, the excess of ammonia being distilled off and titrated with standard acid, calculating the per cent of formaldehyde by the amount of ammonia absorbed.

Detection of Formaldehyde.—Methods have previously been given for the detection of formaldehyde in milk. Pure milk furnishes a convenient reagent for the detection of formaldehyde in various preparations. A solution of the sample to be tested is acidified with phosphoric acid, subjected to distillation, and the first few cubic centimeters of the distillate are tested for formaldehyde as follows:

- (1) Hydrochloric Acid and Ferric Chloride Test.—Add a few drops of the suspected distillate to about 10 cc. of pure milk (previously proved free from formaldehyde) in a porcelain casserole, and carry out the test as described on page 180.
- (2) Hehner's Sulphuric Acid Test.—Apply the test as described on page 180 to 10 cc. of pure milk to which a few drops of the suspected distillate have been added.
- (3) Resorcin or Carbolic Acid Test.—To about 10 cc. of the distillate to be tested, add a few drops of a 1% solution of carbolic acid or resorcin, mix thoroughly, and carefully pour the liquid down the side of a test-tube containing concentrated sulphuric acid. In the presence of formaldehyde, a rose-red zone is formed at the junction of the two liquids, sensitive to 1 part in 200,000. If formaldehyde be present to an extent exceeding 1 part in 100,000, a white turbidity or precipitate is formed above the colored zone.
- (4) Phenylhydrazine Hydrochloride Test.*—One gram of phenylhydrazine hydrochloride and 1½ grams sodium acetate are dissolved in 10 cc. of water. Add 2 to 4 drops of this reagent, and an equal amount of sulphuric acid, to 1 or 2 cc. of the distillate to be tested in a test-tube. A green coloration is produced in the presence of formaldehyde.

^{*} Jour. Am. Chem. Soc., 22, p. 135.

If present in a very small amount (say 1 part formaldehyde in 200,000), heat is necessary to bring out the color.

Determination of Formaldehyde.—The exact quantitative determination of formaldehyde in food products is difficult, owing to its extreme volatility as well as the uncertainty of the compounds which it forms with proteins. A rough idea of the amount present may often be gained by the intensity of the colorations produced in carrying out the various qualitative tests.

Formaldehyde in the small amount present in food products may be roughly determined by the potassium cyanide method (p. 181), on separate portions of the distillate of about 20 cc. each, collecting the distillate as long as an appreciable amount of formaldehyde is shown therein.

BORIC ACID.

Boric or boracic acid is commonly obtained in impure form from lagoons or fumaroles of volcanic origin in Tuscany. It is afterwards purified by recrystallization. It is weakly acid, and readily soluble in water and in alcohol. Its alcoholic solution, even when the acid is present in small quantity, burns with a characteristic green flame. The acid is quite volatile with steam.

Borax, the most commonly known salt of boric acid, is found native in Italy, California, and elsewhere, and is also made from boric acid. It is mildly alkaline, and readily soluble in water.

Boric acid and borax, either used separately or mixed, have long been used as preservatives, especially in animal foods. A mixture of 3 parts boric acid and 1 part borax has been found very effective as a milk and butter preservative, as well as for meat products.

Determination of Boric Anhydride in Commercial Preservatives.— Gladding Method.*—A 150-cc. flask, Fig. 117, is arranged with a doubly perforated stopper having two tubes, one of which, the inlet-tube reaching nearly to the bottom, connects it with a larger flask, while the other or outlet-tube communicates with a Liebig condenser, which in turn delivers into a receiving-flask. In the 150 cc. flask, I gram of the powdered sample is placed, with about 20 cc. of 95% methyl alcohol and 5 cc. of 85% phosphoric acid. The larger flask is then filled two-thirds full of methyl alcohol, and heated on the water-bath after the apparatus has been connected up. Heat is also applied to the 150-cc. flask, the

^{*} Jour. Am. Chem. Soc., 20, 1898, p. 288.

whole arrangement being such that a continuous current of methyl alcohol vapor bubbles through the liquid in the smaller flask, the heat being so regulated that from 15 to 25 cc. of methyl alcohol remains in the 150-cc. flask, while about 100 cc. of distillate passes into the receiving-flask in half an hour. Continue the distillation till all the acid has passed over, which is usually accomplished by distilling 100 cc. By a gentle aspiration upon the receiving-flask, loss by leaking may be avoided.

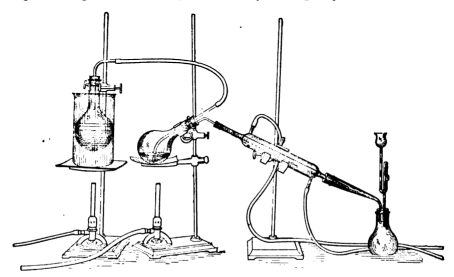


Fig. 117.—Apparatus for Determining Boric Acid According to Gladding.

Prepare a mixture of 40 cc. of glycerin and 100 cc. of water, and carefully neutralize, using phenolphthalein as an indicator. Add this mixture to the distillate, and titrate the whole with tenth-normal sodium hydroxide. Run a blank with the reagents alone, deducting any acidity. For the factors for calculation see page 824.

Detection of Boric Acid and Borates.—These are best tested for in most cases in a solution of the ash of the sample, the quantity to be used for the test depending largely on the case in hand. With meat products and canned goods, about 25 grams are taken for the test, being first made distinctly alkaline with lime water, dried over the water-bath, and burned. The ash is boiled with from 10 to 15 cc. of water, and tests made on the solution. With such products as salt codfish, which is preserved by brushing or coating with boric mixture, portions of the coating may be scraped off and boiled in water, the tests being made on the aqueous solutions.

(1) The Turmeric-paper Test.—The most delicate test for boric acid, free or combined, is made by the aid of turmeric-paper, prepared by soaking a smooth, thin grade of filter-paper in an alcoholic tincture of powdered turmeric. The paper is afterwards dried and cut into strips, which are kept for convenience in a wide-mouthed bottle in a dark place.

Slightly acidulate the ash of the sample to be tested with a few drops of dilute hydrochloric acid, avoiding an excess of acid. Then dissolve the ash in a few drops of water and thoroughly saturate a strip of the turmeric-paper in the solution. On drying the paper, if boric acid either free or combined be present, a cherry-red coloration will be imparted to the paper, the depth of color depending on the amount present. As a confirmatory test, apply a drop of dilute alkali to the reddened paper, and a dark-olive color will be due to boric acid, sharply to be distinguished from the deep-red color produced when an alkaline solution is applied to ordinary turmeric-paper. The turmeric-paper reaction is delicate to 1 part in 8,000.

- (2) Tincture of Turmeric Test.—To the solution to be tested, slightly acidified with hydrochloric acid, add an equal volume of saturated tincture of turmeric in an evaporating-dish, and heat for a minute or two. A red color, light or dark, depending on the amount of the preservative, is produced if boric acid be present, changed to an olive color by the addition of dilute alkali, after cooling.
- (3) The Flame Test.—A few cubic centimeters of alcohol are added to the dish containing the slightly acidulated ash of the sample to be tested, or to the acidulated dried residue from the evaporation of the aqueous solution of the suspected preservative, and after mixing by the aid of a stirring-rod, the alcohol is ignited. In the presence of any considerable portion of free or combined boric acid, a greenish tinge will be observed in the flame of the burning alcohol, especially at the first flash, due to the boric ether formed. This test is by no means as delicate at the paper test.

Determination of Boric Acid in Foods.—(1) Thompson's Method.*—Add 1 or 2 grams of sodium hydroxide to 100 grams of the sample, and evaporate to dryness in a platinum dish. Char the residue thoroughly, and boil with 20 cc. of water, adding hydrochloric acid drop by drop till all but the carbon is dissolved. In burning, avoid too high a heat, simply charring sufficiently to insure a clear solution with water. Transfer by washing to a 100-cc. graduated flask, taking care that the volume does not exceed 50 or 60 cc. Add half a gram of dry calcium chloride, then a few drops

^{*} Analyst, 18, p. 184.

of phenolphthalein solution, and next a 10% solution of sodium hydroxide. till a permanent pink color persists. Finally add 25 cc. of lime-water. By this means all phosphoric acid is precipitated in the form of calcium phosphate. Make up to the 100-cc. mark with water, shake, and pour upon a dry filter. To 50 cc. of the filtrate add sufficient normal sulphuric acid to remove the pink color. Then add a few drops of methyl orange. and continue the addition of sulphuric acid till the vellow is just turned to pink. Tenth-normal sodium hydroxide is then added * till the liquid takes on a faint vellow, excess of alkali being avoided. The salts of the acids present at this time are all neutral to phenolphthalein except boric acid and carbon dioxide. Boil the solution to expel the carbon dioxide, cool, add a little more phenolphthalein, and a quantity of glycerin equal in volume to the solution. Finally titrate with tenth-normal sodium hydroxide to a permanent pink color. Each cubic centimeter of tenthnormal sodium hydroxide equals 0.0062 gram crystallized boric acid, H₂BO₃, or 0.0035 gram boric anhydride, B₂O₃, or 0.00055 gram crystallized borax, Na₂B₄O₇,10H₂O.

(2) Gooch's Method.—Mix 400 to 500 grams of the substance with 10 grams of calcium hydrate, evaporate to dryness over a water-bath in a platinum dish, and burn cautiously to an ash. Dissolve the residue in cold nitric acid, and add an excess of silver nitrate to precipitate the chlorine. Filter, make up to 500 cc. with water, shake, and measure out 25 cc. into a 200-cc. flask fitted with a stopper provided with an outlet-tube, and with a separatory funnel forming virtually a thistle-tube, capable of being closed with a glass stop-cock. Through the outlet-tube connect the flask with a Liebig condenser provided with an adapter which can dip below the liquid in the receiver. As a receiver, use a 150-cc. tared platinum dish, which contains a weighed quantity of ignited lime in water.

Add through the thistle-tube 10 cc. of methyl alcohol to the contents of the flask, close the stop-cock therein, and distill the contents in a paraffin-bath at a temperature of 140° C., constantly stirring the liquid in the receiver to keep it alkaline during the distillation. Add five successive portions of methyl alcohol of 12 cc. each to the distilling-flask, and continue the distillation till all the alcohol has passed over. Finally evaporate to dryness the contents of the platinum dish, and ignite over a blast-lamp to constant weight. Multiply the increased weight due to boric oxide by 2.728 to give the equivalent in borax.

^{*}If the value of the standard alkali solution is not absolutely certain, it had best be restandardized against pure crystallized boric acid, 0.31 gram of which should neutralize 50 cc. of tenth-normal alkali.

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SALICYLIC ACID.

Salicylic acid (HC₇H₅O₃) is a white, crystalline, strongly acid powder, made synthetically by treatment of carbolic acid with sodium hydroxide and carbon dioxide, or naturally from methyl salicylate (which occurs in oil of wintergreen to the extent of about 90%), by treatment of the wintergreen oil with strong potash lye. Most of the commercial salicylic acid is of the synthetic variety. Pure salicylic acid crystallizes from alcoholic solutions in 4-sided prisms, and from aqueous solution in long, slender needles. It melts at 155° to 156° C. It is slightly soluble in cold water (1 part in 450), and much more so in hot water. It is readily soluble in ether, alcohol, and chloroform.

It is frequently found on the market as a food preservative in the form of the much more soluble sodium salt, sodium salicylate, (NaC₇H₆O₅), which is, however, converted into salicylic acid when added to acid-fruit preparations, condiments, and liquors.

Sodium salicylate is a white, amorphous powder, soluble in 0.9 parts water and in 6 parts alcohol. It is prepared by treating salicylic acid with a strong, aqueous solution of sodium carbonate, and afterwards purifying. If a known weight of the powdered preservative be ignited, and a solution of the ash titrated with tenth-normal sulphuric acid, using cochineal as an indicator, each cubic centimeter of the acid is equivalent to 0.0160 gram of sodium salicylate.

Salicylic acid is largely used as a preservative of jellies, jams, and fruit preparations, canned vegetables, ketchups, table sauces, wines, beer, and cider. It is rarely used in milk and milk products, or in meats.

Bucholz has shown that 0.15% of salicylic acid is sufficient to prevent bacteria from developing in ordinary organic substances, while as small a quantity as 0.04% produces a marked restraining influence.

Small amounts of salicylic acid occur naturally in grapes, strawberries, and other fruits, but the amounts are too small to give distinct color reactions when only 50 grams of the fruit products are used for tests.

Detection of Salicylic Acid.—If the sample to be tested is of a similar nature to jelly, jam, ketchup, cider, etc., or capable of getting into aqueous solution, slightly acidify the liquid or pasty material, diluted, if necessary, with weak sulphuric (if not already acid), and shake directly with an equal bulk of ether, petroleum ether, or chloroform, in a corked flask, or in a separatory funnel. If the sample be too thick in consistency to

shake directly, macerate in a mortar with alkaline water, and strain through cloth. Acidify the filtrate with dilute sulphuric acid, and then proceed to shake with the immiscible solvent as above. Separate by decantation or otherwise the immiscible solvent containing the preservative, if present, and allow it to evaporate in an open shallow dish, either at room temperature or at a low heat. In case an emulsion forms on shaking, which is quite apt to happen, especially with ether for a solvent, divide the whole mixture between two tubes of a centrifuge of the form shown in Fig. 11, and whirl for three minutes at a high rate of speed. This usually serves to break up the most obstinate emulsion, so that it is easy to separate by decantation. If a considerable amount of salicylic acid be present, it will sometimes appear in the residue in the form of fibrous crystals.

- (1) To a portion of the dry residue add a drop of ferric chloride solution. A deep purple or violet color indicates salicylic acid.* If doubt exists as to the color, dilute with water, which often serves to bring out a distinctive purple coloration otherwise unobservable.
- (2) Another portion of the residue may be heated with methyl alcohol and sulphuric acid. If salicylic acid be present, the well-known odor of methyl salicylate will be produced.
- (3) A portion of the dry ether extract is warmed gently with a drop of concentrated nitric acid, and two or three drops of ammonia are added. Yellow ammonium picrate will be formed if a considerable quantity of salicylic acid be present, and a thread of wool free from fat may be dyed by soaking therein. This test is by no means as delicate as the ferric chloride color test.

Instead of evaporating the ether solution of the salicylic acid to dryness, the author prefers to shake out the salicylic acid from the ether with dilute ammonia, evaporate the solution of ammonium salicylate nearly to dryness, and apply the tests given above to the concentrated solution. In this case the ether may be recovered.

Determination of Salicylic Acid.—Dubois Method.†—In the case of ketchups and similar pulped materials place 50 grams in a graduated 200-cc. flask, make slightly alkaline with ammonia, add 15 cc. of milk

^{*} Peters (U. S. Dept. of Agric., Bur. of Chem., Bul. 65, p. 160) advises the use of chloroform as more convenient for extraction when testing for salicylic acid, and recommends that the chloroform extract without evaporation be shaken in a test-tube with a drop of ferric chloride reagent and a little water. In the presence of salicylic acid, the violet color will be apparent in the supernatant aqueous layer.

[†] Jour. Am. Chem. Soc., 28, 1906, p. 1616. U. S. Dept. of Agric., Bur. of Chem., Bul. 107 (rev.), p. 179.

of lime (200 grams of quicklime in 2000 cc. water), complete the volume, shake and filter. Transfer 150 cc. of the filtrate to a separatory funnel, acidify with hydrochloric acid, and extract with four portions of 75 to 100 cc. of ether. Wash the combined extract twice with 25 cc. of water, and distil off the ether slowly, allowing the last 20 to 25 cc. to evaporate spontaneously. Dissolve the residue in a small amount of hot water, make up to a definite volume with water, and add to an aliquot portion a few drops of a 2% solution of ferric alum to develop the color. Estimate the amount of salicylic acid by matching the color thus obtained with that produced in a solution containing 1 mg. of salicylic in 50 cc., using either a colorimeter or Nessler tubes for making the comparison.

In the case of semisolid materials, such as mince meat, jams, etc., macerate 50 grams with water in a mortar previous to treatment as above described.

Liquids and solutions of jellies and other materials free from pulp may be extracted with ether directly after acidifying.

BENZOIC ACID.

Benzoic Acid (HC₇H₅O₂) is produced by the oxidation of a large number of organic substances, particularly toluene. It is also extracted by sublimation from gum benzoin, which exudes from the bark of the Styrax benzoin, a tree growing in Java, Sumatra, Borneo, and Siam. Most of the commercial benzoic acid is made from toluene by treatment with chlorine and subsequent oxidation.

Benzoic acid crystallizes in leaflets, having a silky luster. It is odor-less when cold, is soluble in 200 parts of cold, and 25 parts of boiling water, and readily dissolves in alcohol, ether, and chloroform. Its melting-point is 120°, and it sublimes at a slightly higher temperature.

Sodium Benzoate (NaC₇H₅O₂) is the salt most largely used in commercial preservatives, being much more soluble than the acid itself, into which, however, it is converted when put into acid fruit preparations. Sodium benzoate is prepared by adding benzoic acid to a concentrated hot solution of sodium carbonate till there is no longer effervescence, and then cooling, and allowing the sodium benzoate to crystallize out.

In titrating solutions of ignited sodium benzoate with tenth-normal sulphuric acid, each cubic centimeter of the standard acid is equivalent to 0.0144 gram of the benzoate.

Sodium benzoate is a white, amorphous powder, having a sweetish, astringent taste. It is soluble in 1.8 parts of cold water, and in 45 parts of alcohol. Benzoic acid is commonly found as a preservative of ketchups, jellies, jams, and canned goods, and less often in wines and liquors.

Benzoic acid occurs naturally in the cranberry and other berries of the Ericacea.

Detection of Benzoic Acid.—The sample is extracted with ether or chloroform in precisely the same manner as directed for salicylic acid. In fact, it is nearly always desirable to test the same sample for both these preservatives, since either and sometimes both are apt to be found in the same class of food products. For this purpose, the ether or chloroform extract is conveniently divided and evaporated to dryness in separate dishes, one of the residues to be tested for salicylic, and the other for benzoic acid. A considerable amount of benzoic acid is apparent in the residue as shining crystalline scales or needles.

In the author's experience a better procedure than evaporating the ether solution is to extract the benzoic acid from the ether by shaking with ditute ammonia, evaporate the solution of ammonium benzoate nearly to dryness, and apply tests to the concentrated solution.

- (1) Ferric Chloride Test.—A portion of the residue from the ether extract is dissolved in ammonia, and evaporated over the water-bath until neutral to test paper. The residue is stirred in a few drops of warm water, and filtered through a small filter into a narrow test tube. A drop of neutral ferric chloride (prepared by precipitating a portion of the iron from a solution of the salt by ammonia and filtering) is added, and in the presence of benzoic acid a flesh-colored precipitate of ferric benzoate is produced, very characteristic and unmistakable, because of its peculiar color, when the solution in which the test is made is colorless. It occasionally happens, however, in the case of jellies, jams, and ketchups, that these preparations are artificially colored with a dyestuff that persists by its depth of color in obscuring that of the ferric benzoate, especially when only a small amount of benzoic acid is present. Again, in such products as sweet pickles, a precipitate of basic ferric acetate might also come down with the ferric benzoate, and thus confuse. In such cases one of the following methods should be carried out.
 - (2) Sublimation Method.*—Evaporate an ammoniacal solution of the

^{*} Annual Report, Mass. State Board of Health, 1902, p. 486. Food and Drug Reprint, p. 34.

ether extract till neutral in a large watch-glass, by the aid of a gentle heat. Fasten with clips or otherwise a second watch-glass to the first, edge to edge, so as to form a double convex chamber, with a cut filter-paper between. Place upon a small sand-bath and heat. Benzoic acid, if present, will sublime upon the surface of the upper glass in minute needles, recognizable under the microscope. It may further be tested by determining the melting-point of the crystals, or by treating the residue with ammonia, evaporating, and applying the ferric chloride test as above.

- (3) Mohler's Method.*—The ether extract is evaporated and the residue heated with 2 or 3 cc. of strong sulphuric acid till white fumes appear; organic matter is charred and benzoic acid is converted into sulpho-benzoic acid. A few crystals of potassium nitrate are then added. This causes the formation of metadinitrobenzoic acid. When cool, the acid is diluted with water, and ammonia added in excess, followed by a drop or two of fresh, colorless ammonium sulphide. The nitro compound becomes converted into ammonium metadiamidobenzoic acid, which possesses a red color. This reaction takes place immediately, and is seen at the surface of the liquid without stirring.
- (4) Peter's Oxidation Method.†—This method depends on the oxidation of benzoic to salicylic acid by the action of sulphuric acid and barium peroxide, and should, of course, be applied only when salicylic acid has been first proved absent.‡

A portion of the residue, say 0.1 gram, from the ether or chloroform extraction of the suspected sample, is transferred to a large test-tube, and dissolved in from 5 to 8 cc. of concentrated sulphuric acid. Small portions of barium peroxide are then successively added, and the tube shaken in cold water to keep the temperature down, using from 0.5 to 0.8 gram of the peroxide in all. This should produce a permanent froth on the sulphuric acid solution. After standing for half an hour, the test-tube is filled three-quarters full of water, and the mixture shaken, quickly cooled, and filtered. The filtrate is then extracted with ether or chloroform, and the extract tested in the regular manner for salicylic acid.

^{*} Bul. Soc. Chim., 1890, 3, 414; U. S. Dept. of Agric., Bur. of Chem., Bul. 65, p. 109. † U. S. Dept. of Agric., Bur. of Chem., Bul. 65, p. 160.

[‡] In view of the fact that saccharin acts in a similar manner to benzoic acid, the absence of saccharin must also first be established.

Determination of Benzoic Acid.—La Wall and Bradshaw Method.* -Shake thoroughly for five minutes 20 grams of the material, 2 grams of sodium chloride, 5 cc. of hydrochloric acid, and 25 cc. of saturated solution of sodium chloride. Transfer to a moistened filter, collect the filtrate in a graduated 100-cc. flask, and wash the residue on the filter with saturated solution of sodium chloride until the filtrate measures 100 cc. Transfer the filtrate to a separatory funnel and shake out with three portions of chloroform, using 25 cc., 15 cc., and 10 cc. respectively. Evaporate the chloroform at room temperature. If the residue is perfectly white and crystalline, as is usually the case, dry to constant weight over sulphuric acid in a desiccator. If the residue is slightly yellowish and oily, which rarely occurs, dissolve it in about 10 or 15 cc. of weak ammonia, filter into a separatory funnel, washing the filter and funnel with water. Acidulate with dilute sulphuric acid, again shake out with chloroform, evaporate, dry, and weigh.

After obtaining the weight, dissolve the residue in from 3 to 5 cc. of alcohol, and titrate the solution with twentieth-normal potassium hydroxide solution. The volumetric and gravimetric results usually will agree within 1 or 2 milligrams.

Divide the solution after titration, which is slightly alkaline, into two portions. Test one portion for cinnamic acid by adding manganous sulphate solution, which, according to Scoville, forms with cinnamic acid a precipitate, while benzoic acid does not. To the other portion add neutral ferric chloride solution to confirm the presence of benzoic acid.

The foregoing process is based on principles brought to notice by Moerk.† It is not applicable in the presence of salicylic acid or saccharin.

The filtration in the case of high-grade ketchups is often exceedingly slow. This difficulty may be avoided by the following modifications devised by Bigelow: ‡

To 200 grams of ketchup add 20 grams of finely powdered sodium chloride, and enough saturated solution of sodium chloride to make exactly one liter, shake thoroughly and allow to stand over night. Filter through a dry paper, transfer 500 cc. of the filtrate to a separatory funnel, add 5 cc. of sulphuric acid (1:5), and extract successively with 100, 50, 50, and 25 cc. of chloroform. In other respects proceed as in the original method.

^{*}Am. Jour. Pharm., 80, 1908, p. 171. This method, although devised for ketchup, is also applicable to other vegetable products.

[†] Proc. Penn. Pharm. Assn., 1905, p. 181.

[‡] A. O. A. C. Proc., 1908, U. S. Dept. of Agric., Bur. of Chem., Bul. 122, p. 68.

Hilyer's Method.*—This method is valuable as a check on the La Wall and Bradshaw method. After titrating the benzoic acid obtained as described in the preceding section, proceed as follows:

Evaporate to dryness the accurately neutralized solution (which should not have even a slight alkaline reaction), and redissolve in a few cc. of alcohol saturated with silver benzoate. Filter if not clear, wash with a few drops of aldehyde-free alcohol saturated with silver benzoate, and treat with 10 to 15 cc. of a saturated solution of silver nitrate in aldehyde-free alcohol. Collect the precipitate in a Gooch crucible, care being taken that the asbestos filter is so prepared as to afford as rapid a filtration as possible, wash with aldehyde-free alcohol, and finally with a little ether, heat in a water-oven until the ether is removed, cool, and weigh. Care must be taken to perform all the operations as quickly as possible to avoid separation of silver oxide.

The aldehyde-free alcohol is prepared as described on page 745, with the additional precaution of distilling over sodium hydroxide after treatment with metaphenylenediamine hydrochloride.

Sublimation Method.—The ether solution of the benzoic acid obtained, as in the case of salicylic acid, by shaking out with ether (page 826), is evaporated, dried over sulphuric acid, and subjected to sublimation in a suitable form of apparatus. The apparatus shown in Fig. 118, devised by Bird, has been adopted by the A. O. A. C.† The manipulation is as follows:

Transfer the ether solution to tube a, evaporate the ether in a gentle current of air, dry in a vacuum desiccator until the contents are thoroughly dry, and sublime the benzoic acid at 250° C., collecting the sublimate in tube b.

During the sublimation, air is drawn very slowly through the apparatus (a wash bottle is used to gauge the speed of the current) to insure the volatilized benzoic acid passing into tube b. The joint between the two tubes is preferably made by means of a cork stopper. The most satisfactory results are obtained by placing the tube a inside of an oven the temperature of which is raised gradually until it reaches 250° C. The bulb of the tube b should be just outside of the oven, in order that the crystals may form therein. By means of this apparatus considerably higher results are obtained than by subliming on a watch-glass, as described above.

^{*} A. O. A. C. Proc. 1908, U. S. Dept. of Agric., Bur. of Chem., Bul. 122, p. 74-† U. S. Dept. of Agric., Bur. of Chem., Bul. 90, p. 59; Bul, 107 (rev.), p. 181.

The sublimate of benzoic acid collected in tube b may be removed by solution in alcohol, and the amount confirmed by titration. A sublimate is sometimes obtained which somewhat resembles benzoic acid in appearance, and which has an acid reaction. Before applying the method, therefore, to any class of foods, blank experiments should be made to determine whether a sublimate is obtained under the same conditions from the ether extract of that class of foods.

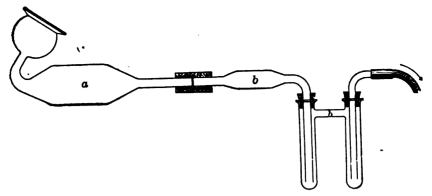


Fig. 118.—Apparatus for Sublimation of Benzoic Acid.

West's Distillation Method.*—1. Apparatus.—The special form of double flask for distillation in a current of steam is the same as that employed by Hortvet† in determining the volatile acids of wine (Fig. 115). The steam tube leading from the outer to the inner flask, being introduced half-way up the side of the inner flask, makes it possible to connect the apparatus in such a way that at the beginning of the operation the water in the outer flask will reach to the height of the contents of the inner flask. The side tube leading from the neck of the outer flask is provided with a rubber tube and pinch-cock for use in relieving the steam pressure and avoiding the danger of drawing the contents of the inner flask over into the outer flask.

2. Process.—Weigh into the inner flask of the apparatus 10 grams, add 1.5 to 2.0 grams of paraffin free from volatile matter, and connect with the condenser. Add 10 cc. of concentrated sulphuric acid, drop by drop, through the funnel tube at such a rate as to complete the addition

^{*} Jour. Ind. Eng. Chem., 1, 1909, p. 190.

[†] Ibid., 1, 1909, p. 31.

in two to three minutes, mix thoroughly by gentle agitation, and allow to stand five to ten minutes after all apparent action of the sulphuric acid has stopped. Measure 150 cc. of distilled water into the outer flask, heat the water slowly to boiling, and continue the boiling until 100 cc. of distillate have been collected, the rate of distillation being such as to yield this amount in 25 to 30 minutes.

Filter the distillate into a separatory funnel, and rinse receiver and filter with two 10-cc. portions of water. Shake with three portions of ether, using 50 cc., 30 cc., and 20 cc., and wash the combined ether extracts by shaking with four 50-cc. portions of water and a last portion of 25 cc., which portion should not require more than a drop of tenth-normal alkali for neutralization, indicating the complete removal of volatile acids. Transfer the ether extract to a tared, wide-mouthed flask, and distil off the ether on the water-bath as quickly as possible. At just the point where ebullition of the ether ceases, remove the flask from the bath, blow air into it to remove the last traces of ether, and dry in a desiccator over night, or until constant weight is secured.

The benzoic acid may also be determined by titration, in which case the filtration of the distillate, also the drying and weighing of the acid, may be omitted. The crystals of benzoic acid are dissolved in alcohol carefully neutralized immediately before each analysis, and the solution titrated with tenth-normal alkali.

SULPHUROUS ACID AND THE SULPHITES.

Free sulphurous acid in the form of sulphur fumes is extensively employed to bleach molasses, to disinfect wine casks, and to bleach and preserve dried fruits. This process is known as "sulphuring." It is stated that the sulphur dioxide combines with the acetaldehyde of wines forming aldehyde-sulphurous acid, which is comparatively harmless. In the case of dried fruits it is believed to form compounds with the sugars.

The sulphurous acid salts most commonly employed as food preservatives are the bisulphites of sodium and calcium, NaHSO₃ and Ca(HSO₃)₂. Others used to some extent are the normal sodium sulphite, and also potassium and ammonium sulphite. The sulphites are usually commercially prepared by passing sulphurous acid gas through strong solutions of the carbonates. Acid sulphites are formed by an excess of the sulphurous acid in the solution of the sulphite. The acid sulphites are distinguishable from the sulphites by their reaction with

litmus paper, the former being acid, while the latter are neutral or feebly alkaline. All of these salts have a bitter, salty, and highly sulphurous taste, and possess a very pungent, irritating odor. With the exception of normal calcium sulphite, all of the above are readily soluble in water.

The sulphites are most commonly used as preservatives of fruit juices, ketchups, fruit and vegetable pulps, wines, malt liquors and meat products. They are frequently mixed with other antiseptics, as with the salts of salicylic and benzoic acids.

Detection and Determination of Sulphurous Acid.—The same methods are used for the detection of sulphurous acid as for its quantitative determination, except that in the former case weighed quantities need not be employed, and the precipitate obtained by the barium sulphate method need not be weighed.

Distillation Method.—This method is adapted to all food products whether solid or liquid.

Place 50 to 200 grams of the material in a 50-cc. flask, add water, if necessary, and 5 cc. of a 20% solution of phosphoric acid, and distil in a current of carbonic acid or steam * into water containing a few drops of bromine, until 100 cc. have passed over. If sulphides are present, as is true of decomposed meat products and possibly other food products, the steam from the distilling-flask before entering the condenser should be passed through a flask containing 40 cc. of a 2% neutral solution of cadmium chloride† or a 1% solution of copper sulphate.‡ These solutions effectually remove the hydrogen sulphide generated by the action of the phosphoric acid, without retaining any appreciable amount of sulphurous acid. To avoid escape of sulphurous acid the condenser tube should dip below the surface of the bromine solution.

After the distillation is complete, boil until the excess of bromine is removed, add drop by drop while boiling an excess of barium chloride, allow to stand over night, filter (preferably in a Gooch crucible with a compact mat of woolly asbestos), ignite, and weigh. From the barium sulphate thus obtained, calculate the amount of sulphur dioxide.

If desired, the distillate may be collected in a receiver containing a measured amount of iodine solution, and the excess titrated with thiosulphate solution, using 1% starch paste as an indicator. This method, however, has the disadvantage that certain volatile organic substances.

^{*} Gudeman, Jour. Ind. Eng. Chem., 1, 1909, p. 81.

[†] Horne, U. S. Dept. of Agric., Bur. of Chem., Bul. 105, p. 125.

[‡] Winton and Bailey, Jour. Am. Chem. Soc., 29, 1907, p. 1499.

act on the iodine solution. After the titration, therefore, add to the solution a few drops of barium chloride reagent. If no appreciable precipitate occurs, the presence of sulphurous acid should be disregarded.

Direct Titration Method.*—This method is applicable to sauternes and other white wines and to beer, but should not be used for other materials, unless found by experiment to yield accurate results.

To 25 grams of the sample, finely divided in water if solid or semi-solid, add 25 cc. of a normal solution of potassium hydroxide in a 200-cc. flask. Shake thoroughly, and set aside for at least fifteen minutes with occasional shaking. 10 cc. of sulphuric acid (1:3) are then added with a little starch solution, and the mixture is titrated with N/50 iodine solution, introducing the iodine solution quite rapidly, and adding it till a distinct fixed blue color is produced. 1 cc. of the iodine solution is the equivalent of 0.00064 gram SO₂.

FLUORIDES, FLUOSILICATES, AND FLUOBORATES.

These substances all possess strong antiseptic qualities, and while no instances are recorded of the use of the last two classes of compounds in this country, the use of fluorides as a preservative of beer is practiced to some extent. The salt most commonly used is ammonium fluoride (NH₄F), preparations of this salt being sold commercially under various trade names as beer preservatives. Ammonium fluoride exists as small, deliquescent, hexagonal, flat crystals. Its taste is strongly saline. It is soluble in water, and slightly soluble in alcohol. Sodium fluoride (NaF) occurs as clear, lustrous crystals, soluble in water.

Detection of Fluorides.—Modification of Blarez' Method.†—Thoroughly mix the sample and heat 150 cc. to boiling. Add to the boiling liquid 5 cc. of a 10% solution of barium acetate. Collect the precipitate in a compact mass, using to advantage a centrifuge, wash upon a small filter, and dry in the oven. Transfer to a platinum crucible, first breaking up the dry precipitate and then adding the filter ash to the crucible. Prepare a glass plate (preferably of the thin variety commonly used for lantern-slide covers) as follows: First thoroughly clean and polish, and coat on one side by carefully dipping while hot in a mixture of equal parts of Canauba wax and paraffin. Near the middle of the plate make a small cross or other distinctive mark through the wax with a sharp

^{*} U. S. Dept. of Agric., Bur. of Chem., Bul. 65, p. 90.

[†] Mass. State Board of Health An. Rep., 1905, p. 498. Chem. News, 91, 1905, p. 39.

instrument, such as a pointed piece of wood or ivory, which will remove the wax and expose the glass without scratching the latter. Add a few drops of concentrated sulphuric acid to the residue in the crucible, and cover with the waxed plate, having the mark nearly over the center, and making sure that the crucible is firmly imbedded in the wax. Place in close contact with the top or unwaxed surface of the plate a cooling device, consisting of a glass cylinder the bottom of which is closed with a thin sheet of pure rubber. Keep the cylinder filled with ice water, so that the wax does not melt. Heat the bottom of the crucible gently over a low flame or on an electric stove for an hour. Remove the glass plate and indicate the location of the distinguishing mark on the unwaxed surface of the plate by means of gummed strips of paper, melt off the wax by heat or a jet of steam, and thoroughly clean the glass with a soft cloth. A distinct etching will be apparent on the glass where it was exposed, if fluoride be present.

Detection of Fluoborates and Fluosilicates.*—Two hundred cc. of the wine or other sample are made alkaline with lime water, evaporated to dryness, and ignited. The crude ash is first extracted with water acidified with acetic acid, and the solution filtered. The insoluble residue is again ignited and extracted with dilute acetic acid, which is filtered off and added to the first extract. The filtrate contains the boric acid, if present, and this is tested for as directed on page 823. Calcium silicate or fluoride, if present, is in the insoluble portion.

Incinerate the filter with the insoluble portion, transfer the ash to a test-tube, mix with some silica, and add a little concentrated sulphuric acid. A small U-tube should be attached to the test-tube, containing a very little water. The test-tube is immersed for half an hour in a beaker of water kept hot on a steam-bath. In the presence of fluoride, silicon fluoride will be generated, and will be decomposed by the water, forming a gelatinous deposit on the walls of the tube.

If both boric and hydrofluoric acids are found, the compound present is undoubtedly a borofluoride. If no boric acid is found, but silicon fluoride is detected, repeat the operation, but without the added silica. If the silicon skeleton is then formed, fluosilicate is probably present.

^{*} U. S. Dept. of Agric., Bur. of Chem., Bul. 59, p. 63.

BETA-NAPHTHOL.

Beta-naphthol ($C_{10}H_7OH$) is a phenol, occurring naturally in coaltar, but the commercial product is more commonly prepared artificially from naphthalene by digesting the latter with sulphuric acid, and fusing the product with alkali. It is a colorless, or pale buff-colored powder, with a faint phenolic odor and a sharp taste. It is slightly soluble in water, and readily soluble in alcohol, ether, and chloroform. Its melting-point is 122° C. In alcoholic solution it is neutral to litmus.

It is used to some extent in alcoholic solution as a preservative of cider.

Detection of Beta-Naphthol.—Bube * states that if an ethereal extract of beta-naphthol is evaporated to dryness, and the residue dissolved in hot water made first faintly alkaline with ammonia, and then faintly acid with very dilute nitric acid, a beautiful rose color will be developed on the addition of a drop of furning nitric acid or of a nitrite. He declares the test to be a delicate one, but it is apparently sometimes obscured by interfering substances, which the ether may dissolve. It should also be carried out in a faint light, as strong sunlight affects the color.

Ferric chloride, when applied to an aqueous solution of beta-naphthol, produces a greenish coloration.

Shake about 50 grams of the sample to be tested with chloroform in a separatory funnel, evaporate the chloroform extract to a small volume (say 1 or 2 cc.), transfer to a test-tube, add 5 cc. of an aqueous solution of potassium hydroxide (1:4), and warm gently. If beta-naphthol is present, a deep-blue color will appear in the aqueous layer, turning through green to light brown.

ASAPROL, OR ABRASTOL.

These are trade names for calcium α -mono-sulphonate of betanaphthol, $Ca(C_{10}H_0SO_3OH)_2$, a white, odorless, scaly powder, sometimes slightly reddish, obtained by the action of heated sulphuric acid on betanaphthol, the resulting compound being afterwards treated with a calcium salt. It is readily soluble in water and alcohol, and is neutral in reaction. Its taste is at first slightly bitter, but rapidly changes to sweet. It decomposes at about 50° C.

^{*} Analyst, 13 (1888), p. 52.

The writer is unaware of any instance of the presence of this substance in foods, but its character is such as to adapt it for use as a preservative of wines and possibly other food products. It has long been regarded as a possible preservative, and the analyst should be prepared to encounter it at any time.

Detection of Asaprol.—Sinabaldi's Method.*—The portion of the solution to be tested (say 50 cc.) is made slightly alkaline with ammonia. and shaken with 10 cc. of amyl alcohol in a separatory funnel. Alcohol is often useful in breaking up an emulsion if there is one. Separate the amyl alcohol extract, which if turbid is filtered, and evaporate to dryness. Wet the residue with about 2 cc. of nitric acid (1:1), heat on the water-bath till the volume is about 1 cc., and wash with a few drops of water into a narrow test-tube. Next add about 0.2 gram of ferrous sulphate and ammonia in excess, a drop at a time, constantly shaking the solution. If a reddish-colored precipitate is formed, it is dissolved by the addition of a little sulphuric acid, and further additions of ferrous sulphate and ammonia are made as before. When a dark-colored or green precipitate appears, add 5 cc. of alcohol, dissolve in sulphuric acid, shake, and filter. If abrastrol be present to the extent of 0.01 gram or more, a red coloration is observed, while in its absence, the filtrate is colorless or faintly yellow.

If the solution to be tested is a fat, it should be melted and extracted with hot 20% alcohol, which is evaporated to dryness, and the above test carried out on the dry residue.

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^{*} Mon. Soi., 1703, (4), 7, p. 842; U. S. Dept. Agric., Bur. of Chem., Bul. 50, p. 01.

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CHAPTER XIX.

ARTIFICIAL SWEETENERS.

UNDER this head are included the intensely sweet coal-tar derivatives, such as saccharin, dulcin, and glucin, that possess no food value whatever in themselves. From their high sweetening power, in some cases several hundred times that of cane sugar, they are capable, when used in minute quantity, of imparting an appropriate degree of sweetness to food products, which, on account of the use of inferior materials, or by reason of the presence of inert or less sweet adulterants, would otherwise be lacking in this property.

Such canned vegetables as sweet corn and peas are subject to treatment with saccharin, especially if by their age and condition before canning they are wanting in the sweet, succulent taste inherent in the fresh product.

The sweetening power of commercial glucose is considerably less than that of cane sugar, so that when large admixtures of the glucose are used in such products as jellies, jams, honey, molasses, maple syrup, etc., to the exclusion of cane sugar, the presence of the glucose might in some cases be suggested by the bland taste of the food, unless reinforced by one of the artificial sweeteners.

The analyst should therefore be on the outlook for one or another of these concentrated sweetening agents in all of the above classes of foods, especially in saccharine products wherein glucose is found to predominate largely over the cane sugar, while the taste is not lacking in sweetness. It is doubtful how far the presence of artificial sweeteners can be regarded as a form of adulteration, unless their presence is legally and specifically prohibited.

SACCHARIN.

Saccharin or Gluside, Benzoyl sulphimide (C₆H₄.CO.SO₂NH), is a white powder, composed of irregular crystals, whose melting-point, when

pure, is about 224° C. It is prepared from toluene, which by treatment with concentrated sulphuric acid is first converted into a mixture of ortho- and para-toluene sulphonic acids. These are further converted into corresponding chlorides, and from the orthochloride, by treatment with ammonia, the imide is formed. It is soluble in 230 parts of cold water, 30 parts of alcohol, and 3 parts of ether. It is sparingly soluble in chloroform, but readily soluble in dilute ammonia. It is from 300 to 500 times as sweet as cane sugar, and, unlike cane sugar, it is not, when pure, charred by the action of concentrated sulphuric acid even on heating. Its aqueous solution is distinctly acid in reaction. Pure saccharin, when heated under diminished pressure, can be sublimed without decomposition.

The addition of 1 part of saccharin to 1,000 parts of commercial glucose renders the latter as sweet as cane sugar.

A sodium salt of saccharin is found on the market, prepared by neutralizing a solution of saccharin with sodium hydroxide or carbonate. The sodium salt crystallizes in the form of rhombic plates, forming a white powder readily soluble in water, and possessing nearly the same sweetening power as saccharin. It is sometimes put up in the form of tablets for the use of diabetic patients as a substitute for sugar.

Saccharin, aside from its sweet taste possesses, according to Fahlberg and List,* antiseptic properties, and on this account it is sometimes used in beer and other liquors. Squibb states that saccharin has about the same power as boric acid as an antiferment

Detection of Saccharin in Foods.—If the sample to be tested is a solution or syrup, render it acid, if not already such, with phosphoric acid, and extract with ether. In case of canned vegetables and similar goods, finely divide the material by pulping or maceration in a mortar, dilute with water, and strain through muslin. Acidify the filtrate, and extract with ether.† If an emulsion forms, use a centrifugal machine (p. 25). Separate the extract, evaporate off the ether, and test the residue for saccharin as follows:

(1) Add to the residue, if it tastes sweet, a few cubic centimeters of hot water, or preferably a very dilute solution of sodium carbonate, in which saccharin is more soluble. An intensely sweet taste is indicative of its presence. This test, if applied directly, will sometimes fail, especially in the case of beer, by reason of the extraction by the ether of various

^{*} Jour. Soc. Chem. Ind., IV, p. 608.

[†] Allen states that a purer residue is obtained if the sample of beer be treated with lead acetate, and filtered before extraction with ether.

bitter principles, such as hop resins, which by their strong, bitter taste mask the sweet taste of saccharin in the residue. Spaeth * recommends that such bitter substances be removed before extraction, which is done by treatment of 500 cc. of the beer with a few crystals of copper nitrate, or with a solution of copper sulphate. The flocculent precipitate formed need not be filtered off, but the liquid is preferably concentrated by evaporation to syrupy consistency, acidified with phosphoric acid, and extracted with three successive portions of a mixture of ether and petroleum ether. After extraction, separation, and evaporation of the solvent, dissolve the residue in weak sodium carbonate. As small a quantity as 0.001% of saccharin can be detected in the final alkaline solution by its sweet taste.

- (2) Bornstein's Test.†—Heat the residue from the ether extraction of the acidified sample with resorcin and a few drops of sulphuric acid in a test-tube till it begins to swell up. Remove from the flame, and, after cooling till the action quiets down, again heat, repeating the heating and cooling several times. Finally cool, dilute with water, and neutralize with sodium hydroxide. A red-green fluorescence indicates saccharin. Gantter ‡ states that it is useless to apply this test to beer, in view of the fact that ordinary hop resin gives the same fluorescence.
- (3) Schmidt's Test.§—The residue is heated in a porcelain dish with about a gram of sodium hydroxide || for half an hour at a temperature of 250° C., either in an air-oven or in a linseed oil bath. This converts the saccharin if present into sodium salicylate. Dissolve the fused mass in water, acidify, and extract the solution with ether. Test the ether residue in the regular manner for salicylic acid with ferric chloride (p. 825), This test can obviously be applied only in the absence of salicylic acid, which should first be directly tested for.

It is recommended that a mixture of equal parts of ether and petroleumether is preferable to the use of ether alone as a solvent of saccharin, as such a mixture, while readily dissolving saccharin, does not, like ether, dissolve other substances, which might form salicylic acid when fused with sodium hydroxide.

Determination of Saccharin.—When saccharin is fused with an alkali and potassium nitrate, the sulphur is oxidized to sulphuric acid. On

^{*} Zeits. angewandte Chem., 1893, p. 579.

[†] Zeits. anal. Chem., 27, p. 165.

[‡] Ibid., 32, 309.

Rep. Anal. Chem., 30; Abs. Analyst, 12, p. 200.

Potassium hydroxide cannot be used instead of sodium hydroxide for the fusion.

this principle depends the following method of Reischauer:* A known quantity of the beer or other liquid to be tested is concentrated by evaporation to about one-third its original volume, acidified with phosphoric acid, and extracted by repeated portions of ether. The combined ether extract is evaporated to small volume, and transferred to a platinum crucible, in which it is further brought to dryness. It is then cautiously ignited with a mixture of about 6 parts sodium carbonate and 1 part potassium nitrate. Dissolve the fusion in water, acidulate with hydrochloric acid, and determine the sulphuric acid in the usual manner with barium chloride. The weight of the precipitated barium sulphate, multiplied by 0.785, gives the weight of saccharin. In view of the fact that only small quantities of saccharin are used in beer and other foods, it is best to employ a large portion of the sample for analysis.

DULCIN.

Dulcin or sucrol, para-phenetol carbamide (C₂H₆O.C₆H₄.NH.CO.NH₂) is a white powder, composed of needle-like crystals, sparingly soluble in cold water, ether, petroleum ether, and chloroform. It dissolves in 800 parts of cold water, 50 parts of boiling water, and 25 parts of 95% alcohol. It is readily soluble in acetic ether. Its melting-point is about 173° C. It is not readily sublimed without decomposition. Dulcin is about four hundred times sweeter than cane sugar.

When a mixture of dulcin and dilute sodium hydroxide is subjected to distillation, phenetidin goes over with the steam into the distillate. When this is heated with glacial acetic acid, phenacetin is formed, which may be tested for as follows: Boil with hydrochloric acid, dilute with water, cool, filter if turbid, and add a few drops of a solution of chromic acid. A deep-red color indicates phenacetin.

Detection of Dulcin in Foods.—In view of the comparatively slight solubility of dulcin in ether and chloroform, acetic ether is the best solvent for purposes of removing it from foods, first making it alkaline.

(1) Bellier's Method.†—A portion of the sample to be tested is made alkaline and extracted with acetic ether. In the case of certain products it is best to subject them to varied preliminary treatment, depending on the case in hand. With such products as thin fruit syrups, simply make alkaline and shake out with acetic ether. In the case of thick fruit syrups, confectionery, and preserves, dilute with water, add an excess of basic

^{*} Abst. Analyst, 11, p. 234.

[†] Ann. de Chim. Anal., 1900, V, pp. 333-337; Abs. Analyst, 26, p. 43.

lead acetate, remove the lead by precipitation with sodium sulphate, filter, and make the filtrate alkaline.

With wines, add 2 grams of mercuric acetate and a slight excess of ammonia, shake, and filter.

With beer, add to 200 cc. 2 or 3 grams of powdered sodium phosphotungstate, and a few drops of sulphuric acid, shake, allow to stand for a few minutes, and filter. Make the filtrate alkaline with ammonia.

Having thus obtained a clarified solution, use from 50 to 200 cc. of neutral acetic ether to say 500 cc. of the alkaline solution, and shake in a separatory funnel. Separate the extract, filter, and evaporate to dryness. If the dulcin exceeds 0.04 gram per liter, crystals will be apparent in the residue. If fats and resins are present in the residue, make repeated extractions with hot water, and evaporate to dryness. The purified residue is finally brought to dryness in a porcelain dish, and treated with 1 or 2 cc. of sulphuric acid and a few drops of a solution of formaldehyde. Let it stand for fifteen minutes, and afterwards dilute with 5 cc. of water. A turbidity or precipitate indicates dulcin.

- (2) Jorissen's Test.*—The residue from the acetic ether extract of an alkaline solution of the sample is treated with 2 or 3 cc. of boiling water in a test-tube, and a few drops of mercuric nitrate † are added. Heat the tube and its contents for five minutes in a boiling water-bath, withdraw, and disregarding any precipitate, add a small quantity of lead peroxide. On the subsidence of the precipitate, which quickly occurs, a fine violet color appears for a short time in the clear upper layer in presence of 0.001 gram of dulcin.
- (3) Morpurgo's Method.‡—To the acetic ether residue, evaporated to dryness in a porcelain dish, add a few drops of phenol and concentrated sulphuric acid, and heat a few minutes on the water-bath. After cooling, transfer to a test-tube, and with the least possible mixing pour ammonia or sodium hydroxide over the surface. A blue zone at the plane of contact between the two layers indicates dulcin.

Determination of Dulcin.—For a quantitative determination, Bellier's method is carried out on a weighed or measured portion of the sample, as follows: In the case of alcoholic beverages first expel the alcohol by

^{*} Chem. Zeit. Rep., 1896, p. 114.

[†] The mercuric nitrate is prepared by dissolving 2 grams of mercuric oxide in dilute nitric acid, adding sodium hydroxide solution till a slight permanent precipitate is formed, diluting to 15 cc., and decanting the clear liquid.

[‡] Zeits. anal. Chem., 1896, 35, p. 104; U. S. Dept. of Agric., Bur. of Chem., Bul. 65, p. 89.

evaporation, and make up to the original volume with water. Treat the various food preparations with the appropriate clarifying reagents, as in Bellier's qualitative test (p. 845), and, after filtering and making alkaline, extract twice with 50 cc. each of acetic ether. The residue is purified if necessary by extraction with hot water as above described, and the final residue is dissolved in 1 to 5 cc. of concentrated sulphuric acid. A few drops of formaldehyde are added. The solution is allowed to stand for fifteen minutes, and then diluted to ten times its volume with distilled water. After twenty-four hours, collect the precipitate on a tared filter, wash with water, dry, and weigh.

GLUCIN.

This comparatively new sweetening agent is the sodium salt of a mixture of the mono- and di-sulphonic acids of a substance having the composition C₁₉H₁₆N₄. In the market it appears as a light-brown powder, readily soluble in water. It is insoluble in ether and chloroform. It decomposes without melting at about 250° C. It is three hundred times sweeter than cane sugar.

A color reaction with glucin is obtained by dissolving it in dilute hydrochloric acid, cooling by immersing the test-tube in water, and to the cold solution adding a little sodium nitrite solution. Finally, to the liquid is added a few drops of an alkaline solution of beta-naphthol, and a red coloration is produced. With resorcin or salicylic acid in alkaline solution, the color will be yellow.

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CHAPTER XX.

FLAVORING EXTRACTS AND THEIR SUBSTITUTES.

Or the three great groups of organic compounds essential for nutrition, the fats and proteins in a state of purity are almost tasteless, as is also true of starch, dextrin, and cellulose of the carbohydrate group. Only the sugars have a pronounced taste. The flavor of food products, aside from their sweetness, is largely due to minor constituents, such as organic acids, ethers, essential oils, etc., which serve chiefly to render the products acceptable to the palate, thereby contributing to their digestibility. Many culinary preparations lacking in flavor, but not in nutritive value, are commonly mixed with substances which supply this deficiency. Spices and flavoring extracts belong to the class of materials added mainly if not entirely for their zest-giving properties.

By far the most extensively used flavoring extracts are those of vanilla and lemon, and in comparison with these the sale of all other varieties is comparatively insignificant. These two favorite extracts are employed in nearly every household, and form a necessary adjunct to almost all forms of desserts, cakes, and confections, as well as to a wide variety of commercial preparations. Others of some importance are extracts of orange, almond, wintergreen, peppermint, rose, and certain spices. Imitation fruit flavors are used in cheap confectionery, ice cream, etc., and are of questionable wholesomeness.

VANILLA EXTRACT.

The Vanilla Bean is the source of pure vanilla extract, besides being used in chopped form directly as a flavoring agent. It is the fruit of the plant of the Vanilla planifolia, or flat-leaved vanilla. This climbing, perennial plant belongs to the orchid family, and is indigenous to Central and South America and the West Indies, but by far the highest prized beans are cultivated in Mexico. While different varieties differ in some details, the best cured beans of commerce, as a rule, are from 20 to 25 cm. in length and from 4 to 8 mm. thick, drawn out at their ends and curved

at the base. They are rich dark brown in color, of a soapy or waxy nature to the touch, deeply rifted lengthwise, and covered with fine frost-like crystals of vanillin. When cut cross-wise, the bean exudes a thick, odorless juice, containing calcium oxalate crystals.

The cross-section of the bean is ellipsoidal in shape. The thick brown walls inclose a triangular cavity, in which are the lobed placentas. Between these are papillæ, secreting a finely granular, yellow, balsamlike substance that contributes much to the flavor of the extract, and helps to give the cut bean its delicious odor.

When first gathered, the beans are yellowish green, fleshy, and without odor, developing their peculiar consistency, color, and smell by the process of fermentation or "sweating," which differs in various countries. According to the best methods the beans are sun-dried for nearly a month, being alternately pressed lightly between the folds of blankets, and exposed to the air. After the curing, they are packed in bundles.

Quicker methods of curing consist of the use of artificial heat and calcium chloride for drying, but the products so prepared are considered inferior in quality.

The Mexican vanilla beans are of the choicest grade, and command a high price, sometimes reaching fifteen dollars per pound. The Bourbon beans, grown in the Isle of Réunion, are next in grade. These beans are shorter than the Mexican and much less expensive. They resemble the Tonka bean in odor. Beans from Seychelles and Mauritius are even shorter than the Bourbon beans, and are largely exported to England. Cheaper varieties are those from South America, which do not bring half the price of the Mexican beans, and the cheapest are the Tahiti beans and so-called "vanillons," or beans of the wild vanilla (Vanilla pompona). These latter are used more in sachet powders and perfumes, possessing an odor not unlike heliotrope.

Composition of the Vanilla Bean.—The following are results of the analyses of two varieties of vanilla beans, according to König:

	A.	В.
Water	25.85	30.94
Nitrogen bodies	4.87	2.56
Fat and wax	6.74	4.68
Reducing sugar	7.07	9.12
Non-nitrogen substances	30.50	32.90
Cellulose	19.60	15.27
Ash	4.73	4.53

Vanillin.—Under "non-nitrogen substances" in the above table is included vanillin, the principle to which vanilla owes its peculiar odor. This body (C₈H₈O₃) is the methyl ether of protocatechuic aldehyde, and is found on the surface of the bean in fine crystalline needles. It has a sharp but pleasant flavor, is soluble with difficulty in cold water, but readily soluble in hot water, ether, alcohol, and chloroform. Its meltingpoint is 80° to 81° C. and it sublimes at 280°. It is present in vanilla beans to an amount varying from 1 to 2½ per cent, and it is a curious fact that varieties of bean most highly prized possess the least vanillin. This is shown by Tiemann and Harmann as follows:

Mexican beans	1.69%	vanillin
Bourbon beans	2.48%	"
Java beans	2.75%	"

While vanillin may be readily extracted by alcohol and other solvents from the beans, such a product would be far too expensive to compete with the commercial synthetic vanillin, an artificial product, chemically identical with the vanillin from the bean. Synthetic vanillin was formerly made from the glucoside coniferin by oxidation with chromic acid. It is now largely obtained by oxidizing the eugenol of clove oil with alkaline potassium permanganate.

If ferric chloride be added to an aqueous solution containing vanillin, a dark-blue coloration will be produced.

Besides vanillin, the bean contains notable quantities of wax, fat, sugar, tannin, gum, and resin.

Exhausted Vanilla Beans are sometimes found on sale, which have been deprived of their vanillin by being soaked in alcohol, after which they are coated with some artificial substitute, presenting the same frosty appearance as the natural vanillin crystals. This may be accomplished by rolling the beans in benzoic acid. Benzoic acid crystals are readily distinguished from those of vanillin under the microscope.

Composition of Vanilla Extract.—Vanilla extract is a dilute alcoholic tincture of the vanilla bean, sweetened by cane sugar. To be perfectly pure it should contain no other added substances, with the possible exception of glycerin, and many of the best brands are free from this. In practice it is variously prepared, but the following method of the U. S. Pharmacopæia is a typical one:

"Vanilla, cut into small pieces and bruised, 100 grams.

[&]quot;Sugar, in coarse powder, 200 grams.

"Alcohol and water, each, a sufficient quantity to make 1,000 cc. "Mix alcohol and water in the proportion of 650 cc. of alcohol to 350 cc. of water. Macerate the vanilla in 500 cc. of this mixture for twelve hours, then drain off the liquid and set it aside. Transfer the vanilla to a mortar, beat it with the sugar into a uniform powder, then pack it in a percolator, and pour upon it the reserved liquid. When this has disappeared from the surface, gradually pour on the menstruum, and continue the percolation, until 1,000 cc. of tincture are obtained."

The best extracts are produced by allowing the cut beans to macerate in the alcohol for several months.

Five vanilla extracts, made by Winton and Silverman from beans of different grades, strictly according to the pharmacopœial formula as above, were analyzed by them with the following results:

ANALYSES OF VANILLA EXTRACTS, U. S. P., MADE IN THE CONNECTICUT EXPERIMENT STATION LABORATORY.

Grade of Bean.	Specific Gravity.	Vanillin. Per Cent.	Alcohol by Weight, Per Cent.	Total Residue, Per Cent.	Cane Sugar, Per Cent.	Residue Other than Cane Sugar, Per Cent.
Mexican (whole). '' (cut)	1.0159 1.0146 1.0109 1.0166 1.0104	0.125 0.065 0.215 0.138	37.96 39.92 38.58 38.32 38.84	22.60 23.10 22.00 23.13 21.75	19.90 19.20 19.00 20.40	2.70 3.90 3.00 2-73 1.75

Vanillin Content.—The writer has found in his examination of a large number of brands of vanilla extract that gave every indication of purity that the content of vanillin varied from 0.05 to 0.200 per cent. It is rare that a pure extract will show more vanillin than the latter figure, though one of Winton's extracts runs as high as 0.215. The writer has found extracts with 0.250 or more of vanillin, but believes them to have been reinforced with the artificial substance. The flavor of the extract depends not only on the vanillin, but also on the various resinous and other extractive matters which it contains.

Use of Alkali.—Some manufacturers employ dilute alkali, generally potassium bicarbonate, to aid in dissolving out the resinous matter from the bean, and to enable them to use a more dilute alcohol. The resulting product made by this process is distinctly inferior, both in taste and odor.

Alcohol in pure extracts varies between the limits of 20 and 40 per cent.

The Tonka Bean forms the basis of many of the cheaper so-called vanilla extracts on the market. It is the seed of the large tree, native to

Guiana, known as *Dipterix* (or *Coumarouna*) odorata. The pods are almond-shaped, and contain a single seed, from 3 to 4 cm. long, shaped like a kidney bean, of a dark-brown color, having a thin, shiny, rough, brittle skin, and containing a two-lobed oily kernel.

Coumarin (C₉H₆O₂), the active principle of the Tonka bean, is the anhydride of coumaric acid. It occurs in the crystalline state between the lobes of the seed kernel. Coumarin occurs also in many other plants. It may be extracted from the beans by treatment with alcohol. It crystallizes in slender, colorless, needles, melting at 67° C. It has a fragrant odor and burning taste. It is very slightly soluble in cold water, but readily soluble in hot water, ether, chloroform, and alcohol. One pound of cut beans yields by alcoholic extraction about 108 grains of coumarin. The latter may be synthetically prepared by heating salicylic aldehyde with sodium acetate and acetic anhydride, forming aceto-coumaric acid, which decomposes into acetic acid and coumarin.

The author has found that an aqueous solution of coumarin, unlike vanillin, forms a precipitate when iodine in potassium iodide is added in excess, the precipitate being at first brown and flocculent, afterwards, on shaking, clotting together to form a dark-green, curdy mass, leaving the liquid perfectly clear.

U. S. Standards.—Vanilla extract is the flavoring extract prepared from the vanilla bean, with or without sugar or glycerin, and contains in 100 cc. the soluble matters from not less than 10 grams of the vanilla bean.

Vanilla bean is the dried, cured fruit of Vanilla planifolia Andrews.

Tonka extract is the flavoring extract prepared from tonka bean, with or without sugar or glycerin, and contains not less than 0.1% by weight of coumarin extracted from the tonka bean, together with a corresponding proportion of the other soluble matters thereof.

Tonka bean is the seed of Coumarouna odorata Aublet (Dipteryx odorata (Aubl.) Willd.).

The Adulteration of Vanilla Extract consists chiefly in the use of coumarin or extract of the Tonka bean, and in the substitution of artificial vanillin, either alone or with coumarin, for the true extractives of the vanilla bean. Imitation vanilla flavors more often consist of a mixture of either tincture of Tonka or coumarin with vanillin in weak alcohol, colored with caramel, or occasionally with coal-tar colors. Or the exhausted marc from high-grade vanilla extract is macerated with hot water and extracted, the extract being reinforced with artificial vanillin or coumarin, or both. A pure vanilla extract possesses

certain peculiarities with regard to its resins and gums that distinguish it from the artificial, or indicate whether or not it has been tampered with. While it is possible to introduce artificial resinous matter in the adulterated brands with a view to deceiving the analyst, it is almost impossible to do this without detection, since different reactions are readily apparent in this case from those of the pure extracts.

Prune juice is said to be used to give body and flavor to vanilla extract. The writer has found spirit of myrcia or bay rum in a sample of alleged vanilla extract, containing also vanillin and coumarin. The adulterant in this sample was present to such an extent as to be unmistakable by reason of the odor.

Factitious Vanilla Extracts are ordinarily indicated (1) by the presence of coumarin, (2) by the peculiar reactions of the resinous matter, or by the entire absence of these resins, (3) by the scanty precipitate with lead acetate, and (4) by the abnormally low or high content of vanillin.

The following figures show the content of vanillin and coumarin in a few typical cheap "vanilla" extracts, selected from a large number examined by the author. All of these were entirely artificial, and ranged from 5 to 20 per cent by weight of alcohol.

	Vanillin, Per Cent.	
A	0.040	0.074
B	None	0.172
C	None	0.330
D	0.250	None
E	0.025	0.144

As a rule these cheap artificial preparations possess considerable body and flavor, but the latter is of a much grosser nature than the genuine vanilla extract, with the delicate and refined flavor of which they are not to be mistaken by any one at all familiar with both varieties.

Winton and Bailey* have found as high as 2.55% of vanillin in imitation extracts. They also have detected the presence of acetanilide in amounts varying up to 0.15%. This substance at one time was extensively employed as an adulterant of vanillin, hence its presence in imitation extracts prepared from such vanillin. It is not only worthless as a flavor, but is a menace to health.

^{*} Conn. Agric. Exp. Sta., Rep. 1905, p. 131.

The author is inclined to place 0.05% of vanillin as a minimum for genuine vanilla extract properly prepared and makes a practice of classing as not of good standard quality those samples that fall below.

METHODS OF ANALYSIS OF VANILLA EXTRACT

Detection of Artificial Extracts.—The presence of coumarin or Tonka tincture to any appreciable extent in vanilla extract is usually recognizable by the odor, to one skilled in examining these flavors. The odor of coumarin is more pungent and penetrating than that of vanillin, and in mixtures is apt to predominate over the milder and more delicate odor of vanillin.

Add normal acetate of lead solution to a suspected extract. The absence of a precipitate is conclusive evidence that it is artificial. If a precipitate is formed, much information may be gained by its character. A pure vanilla extract should yield with lead acetate a heavy precipitate, due to the various extractives. The precipitate should settle in a few minutes, leaving a clear, supernatant, partially decolorized liquid. If only a mere cloudiness is formed, this may be due to the caramel present, and in any event is suspicious.

Examination of the Resins.—Resin is present in vanilla beans to the extent of from 4 to 11 per cent, and the manufacturer of high-grade essences endeavors to extract as much as possible of this in his product. This he can do by the use of 50% alcohol, in which all the resin is readily soluble, or by employing less alcohol and relying on the use of alkali to dissolve it. A pure extract free from alkali should produce a precipitate, when a portion of the original sample is diluted with twice its volume of water and shaken in a test-tube.

When, moreover, the alcohol is removed from such an extract, the excess of resin is naturally precipitated.

The character of the resins extracted from the vanilla bean is so different from that of other resins as to furnish conclusive tests, worked out by Hess * as follows: 25 to 50 cc. of the extract are de-alcoholized by heating in an evaporating-dish on the water-bath to about one-third its volume. Make up to the original volume with water, and, if no alkali has been used in the manufacture of the preparation, the resin will be in the form of a brown, flocculent precipitate. To entirely set free the resin, acidify, after cooling, with dilute hydrochloric acid, and allow to stand till all the resin has settled out, leaving a clear supernatant liquid. The resin may be quantitatively determined, if desired, by filtering, wash-

^{*} Jour. Am. Chem. Soc., 21 (1899), p. 721.

ing, drying, and weighing, but in this case should stand for a long time before filtering.

The resin is collected on a filter, washed, and subjected to various tests. A piece of the filter with the attached resin is placed in a beaker, containing dilute potassium hydroxide. Pure vanilla resin dissolves to a deep-red color, and is reprecipitated on acidifying with hydrochloric acid. Dissolve another portion of the precipitate in alcohol, and divide the alcoholic solution into two portions, to one of which add a few drops of ferric chloride, and to the other hydrochloric acid. Pure vanilla resin shows no marked coloration in either case, but foreign resins nearly all give color reactions under these conditions.

Tannin.—Test a portion of the filtrate from the resin for tannin by the addition of a few drops of a solution of gelatin. A small quantity of tannin only should be indicated, if the extract is pure, a large excess tending to show added tannin.

Determination of Vanillin.—Vanillin may be determined (1) colorimetrically, or (2) by extraction and weighing. The former is by far the quicker and more economical method, since it may be carried out directly in a very small portion of the original alcoholic extract. When, as in some instances, the analyst has only one small bottle of vanilla extract for analysis, it becomes a matter of importance to use as little as possible for each determination. The determination of vanillin by both methods should give concordant results.

Colorimetric Method.*—This is carried out in the author's laboratory as follows: 2 cc. of the vanilla extract are measured into a testtube, and sufficient lead hydrate is added to completely decolorize it. The
mixture is washed upon a filter, and filtrate and washings are collected
in a Nessler tube. Bromine water is then added, after which enough
of a freshly prepared \(\frac{1}{2} \)% ferrous sulphate solution is added to get the
maximum bluish-green color that will be produced, if vanillin is present.

A standard vanillin solution is freshly prepared by dissolving 50 mgm.
of pure vanillin in 25 cc. of alcohol, and making up to 100 cc. with water.
A series of color standards is then made, taking, for instance, \(\frac{1}{2} \), 1, 1\(\frac{1}{2} \), 2\(\frac{1}{2} \), 3, etc., cc. of the vanillin solution in 50 cc. Nessler tubes, each being
treated with two or three drops of bromine water, and with the ferrous
sulphate solution, and made up to the 50-cc. mark.

The lead hydrate is prepared by dissolving 200 grams of lead acetate in 850 cc. of water. The solution is filtered, a solution of potassium

^{*} Massachusetts State Board of Health, An. Rep., 1899, p. 629.

hydroxide is added in excess, and the precipitated hydrate is washed thoroughly several times by decantation, or until neutral. Keep an excess of water in the reagent bottle, and shake up to form a heavy, white emulsion before adding to decolorize. Unless the lead hydrate is washed free from alkali, the latter will precipitate the iron salt when added.

If, for example, 2 cc. of a sample extract, treated as above, are found to give a color corresponding in depth to that produced by 5.5 cc. of the standard solution, the percentage of vanillin would be thus calculated:

```
100 cc. standard solution contain 0.050 gram vanillin.

1 cc. " " 0.0005 " "

5.5 cc. " " 0.00275 " "
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Since 2 cc. of the sample are equivalent to 5.5 cc. of the standard solution, it follows that

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2 cc. of sample contain 0.0275 gram vanillin.
.: 100 cc. " " 0.1375 " "
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In order to avoid calculation of each determination when a large number of extracts have to be examined, the following table will be found useful for expressing results, where the above method of procedure has been exactly carried out:

Number of Cub			
meters of Sta			Equivalent
Vanillin Sol		1	Per Cent of
Correspond			Vanillin
to 2 cc. of Sa	•		in Sample.
0.25	• • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	0.00025
0.5		• • • • • • • • • • • • • • • • • • • •	0.0125
0.75	•••••		0.01875
I.		••••••	0.025
1.5		• • • • • • • • • • • • • • • • • • • •	0.0375
2.		• • • • • • • • • • • • • • • • • • • •	•
2.5	• • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	0.0625
3•	•••••		0.075
3-5			0.0875
4.	•••••		0.1
4.5	•••••		0.1125
5•	• • • • • • • • • • • • • • • • • • • •		0.125
5-5	•••••	••••••	0.1375
6.	•••••	••••••	0.15
7-	• • • • • • • • • • • • • • • • • • • •		0.175
8.	•••••		0.2
9.			0.225
10.	•••••	•••••	0.25

^{* 0.05} gram vanillin in 100 cc.

Determination of Vanillin, Coumarin, and Acetanilide.—Hess and Prescott Method, Modified by Winton and Bailey.*—Weigh 25 grams of the extract into a 200-cc. beaker with marks showing volumes of 25 and 50 cc. Dilute to the 50-cc. mark, and evaporate in a water-bath to 25 cc. at a temperature in the bath of not more than 7c° C. Dilute a second time to 50 cc. and evaporate to 25 cc. Add normal lead acetate solution drop by drop until no more precipitate forms. Stir with a glass rod to facilitate flocculation of the precipitate, filter through a moistened filter, and wash three times with hot water, taking care that the total filtrate does not measure more than 50 cc. Cool the filtrate, and shake with 20 cc. of ether in a separatory funnel. Remove the ether to another separatory funnel, and repeat the shaking of the aqueous liquid with ether three times, using 15 cc. each time. Shake the combined ether solutions four or five times with 2% ammonium hydroxide, using 10 cc. for the first shaking, and 5 cc. for each subsequent shaking. Reserve the combined ammoniacal solutions for determination of vanillin.

Wash the ether solution into a weighed dish, and allow the ether to evaporate at the room temperature. Dry in a desiccator, and weigh. Stir the residue for fifteen minutes with 15 cc. of petroleum ether (boiling point 30 to 40° C.) and decant the clear liquid into a beaker. Repeat the extraction with petroleum ether two or three times. If the residue is completely dissolved by this treatment, the absence of acetanilide and other impurities in the coumarin is assured. Should an appreciable amount of material remain undissolved, allow the dish and contents to stand in the air until apparently dry, completing the drying in a desiccator. Weigh, and deduct the weight from the weight of the residue obtained after the ether evaporation, thus obtaining the weight of the coumarin. The petroleum ether residue, if acetanilide, should melt at about 112° C. and respond to Ritsert's test (p. 859).

Allow the petroleum ether extract to evaporate at room temperature. If it is pure coumarin, it should have a melting-point within a few degrees of 67° C. and respond to the author's test (p. 859).

Slightly acidulate the ammoniacal solution reserved for vanillin with 10% hydrochloric acid. Cool, and shake out in a separatory funnel with four portions of ether, as described for the first ether extraction. Evaporate the ether at room temperature in a weighed dish, dry over sulphuric acid, and weigh.

^{*} Jour. Am. Chem. Soc., 21, 1899, p. 256.; 24, 1902, p. 1128; 27, 1905, p. 719.

If acetanilide has not been previously detected, this residue should be pure vanillin with a melting-point within a few degrees of 80° C.

If acetanilide has been detected, dissolve the residue in 15 cc. of 10% ammonium hydroxide, and shake twice with an equal volume of ether. Evaporate the ether solution at room temperature, dry in a desiccator, and weigh. Deduct this weight from the previous weight, thus obtaining the weight of pure vanillin. The total weight of the acetanilide is obtained by adding the weight of this last extract to that of the residue previously obtained and identified as acetanilide.

In doubtful cases the ammoniacal solution should be acidified, shaken out with ether, and the melting-point of the vanillin, obtained by evaporation at room temperature, determined.

In case the colorimetric method for vanillin was used, and coumarin only is to be separated for gravimetric determination, the author has found that good results are usually obtained by simply treating the dealcoholized original sample with ammonia, extracting it with 3 or 4 portions of chloroform in a separatory funnel, and evaporating the combined chloroform extract in a tared dish at a temperature not exceeding 60° in the oven.

Many of the precautions employed in carrying out the above processes for vanillin and coumarin determination may be dispensed with, if these substances are simply to be tested for qualitatively.

Leach's Test for Coumarin.—The residue, believed to be coumarin, obtained as described in the preceding section, is identified by the following test: Add a few drops of water, warm gently, and add to the solution a little iodine in potassium iodide, reagent No. 143. In presence of coumarin a brown precipitate will form, which, on stirring with the rod, will soon gather in dark-green flecks. The reaction is especially marked if done on a white plate or tile.

Ritsert's Tests for Acetanilide.*—Boil the acetanilide, obtained as described on page 858, in a small beaker for two or three minutes with 2 to 3 cc. of concentrated hydrochloric acid, cool, divide into three portions, and test in small tubes (4 to 5 mm. inside diameter), or by spotting on a porcelain plate, as follows:

(1) To one portion add carefully 1 to 3 drops of a solution of chlorinated lime (1:200) in such a manner that the two solutions do not mix. A beautiful blue color formed at the juncture of the two liquids indicates acetanilide.

^{*} Pharm. Ztg., 33, 1888, p. 383; Abs. Zeits. anal. Chem., 27, 1888, p. 667.

- (2) To another portion add a small drop of potassium permanganate solution. A clear green color is formed if any appreciable amount of acetanilide is present.
- (3) Mix the third portion with a small drop of 3% chromic acid solution. Acetanilide gives a yellow-green solution, changing to dark green on standing five minutes, and forming a dark blue precipitate on addition of a drop of caustic potash solution.

These tests are conclusive only when taken in conjunction with the melting-point.

Vanillin and Coumarin Crystals Under the Microscope.—These substances are best examined when crystallized from ether solution, and several crystallizations may be found necessary, before the best results are obtained. For examination, pour a few drops of the ether solution of the purified vanillin or coumarin directly on a slide, and allow to evaporate spontaneously. Under best conditions vanillin crystallizes from ether in long, slender needles, often radiating from central points, or forming star-shaped bundles.

Coumarin crystals are shorter and thicker than vanillin.

With polarized light pure vanillin crystals give a brilliant play of colors between crossed nicols, even without the selenite plate, while pure coumarin crystals without the selenite are almost lacking in varying colors, and show very little play, even when the selenite is employed. This sharp distinction is not true, when crystallized from chloroform.

Determination of Glycerin.—The presence of any considerable quantity of glycerin is apparent by the character of the residue obtained on evaporating 5 grams to dryness, in the determination of total solids. The residue, if glycerin is present in notable amount, appears of a moist consistency, even when a practically constant weight has been attained at 100° C.

To determine glycerin, proceed as with wines (p. 703).

Determination of Alcohol.—Measure out 25 cc. of the sample, dilute to 50 cc. with water, and distil off about 20 cc. in a 25-cc. graduated receiver. Make up to the mark with water, determine the specific gravity at 15.6°, and find from the alcohol table the per cent corresponding.

Cane Sugar and Glucose are determined as in the case of preserves and jellies.

Detection of Caramel.—Lead Acetate Method.—Dealcoholize, precipitate with lead acetate, and filter, as described for the determination of vanillin, coumarin, and acetanilide (page 858). If the extract is pure,

the filtrate will be light yellow; if colored with caramel, the filtrate will be yellow-brown or deep brown, according to the amount present.

Color Quotient.+Method of Woodman and Newhall.*—Evaporate 35 cc. of the sample, or, if necessary, a diluted portion of the sample, on the water-bath to one-third its volume, and replace the loss by addition of water. Shake the dealcoholized extract in a separatory funnel with three successive portions of 15 cc. each of ethyl acetate. After the last portion has been separated, reserve part of the aqueous layer for comparison, and shake 30 cc., or as much as possible of the remainder with 15 grams of fuller's earth, allow to stand for half an hour and filter. The percentage of color removed by ethyl acetate, divided by the percentage of color removed by fuller's earth, gives the "color quotient." For example, a standard extract gave a color quotient of 2.30, whereas a caramel solution gave 0.276.

Coal-tar Colors are detected by the usual tests (pp. 793 to 812).

LEMON EXTRACTS.

Spirit or essence of lemon of the National Formulary and former editions of the Pharmacopæia, is a 5% solution (by volume) of lemon oil in deodorized alcohol, colored with lemon peel.

This preparation was dropped from the 8th revision of the Pharmacopæia, and *Tinctura limonis corticis* or tincture of lemon peel added. The following are the directions for the preparation of the latter:

- "Macerate the lemon peel in a stoppered, wide-mouthed container, in a moderately warm place, with 1000 cc. of alcohol during forty-eight hours, with frequent agitation; then filter through purified cotton, and, when the liquor has drained off completely, gradually pour on enough alcohol to make 1000 cc. of tincture, and filter."
- U. S. Standards. Lemon Extract is the flavoring extract prepared from oil of lemon, or from lemon peel, or both, and contains not less than 5% by volume of oil of lemon.

Oil of Lemon is the volatile oil obtained, by expression or alcoholic solution, from the fresh peel of the lemon (Citrus limonum L.), has an

^{*} Tech. Quar., 21, 1908, p. 280.

optical rotation (25° C.) of not less than $+60^{\circ}$ in a 100-mm. tube, and contains not less than 4% by weight of citral.

Terpeneless Extract of Lemon is the flavoring extract prepared by shaking oil of lemon with dilute alcohol, or by dissolving terpeneless oil of lemon in dilute alcohol, and contains not less than 0.2% by weight of citral derived from oil of lemon.

Terpeneless Oil of Lemon is oil of lemon from which all or nearly all of the terpenes have been removed.

The U. S. standard for lemon extract (5% of lemon oil by volume) is a fair one. In fact there are commercial extracts on the market containing as high as 12%. An extract of lemon to contain 5% of lemon oil must contain at least 80% by volume of alcohol, lemon oil being insoluble in dilute alcohol. Deodorized, or purified alcohol, commonly known as cologne spirits or perfumers' alcohol, is used in the highest-grade preparations, since the odor of ordinary commercial alcohol produces a slightly deleterious effect.

Adulteration of Lemon Extracts.—For making a cheap extract the cost of the lemon oil is not so important an item as that of the alcohol, and as little as possible of the latter is employed, though pure oil is doubtless used. These terpeneless extracts are made by rubbing the oil in carbonate of magnesia in a mortar, stirring in slowly a little strong alcohol, and allowing the mixture to soak for some time. A varying amount of water is added a little at a time, and the whole is shaken and again allowed to stand, sometimes for a week, before filtering. Finally the extract is filtered, and the coloring matter added, consisting sometimes of turmeric tincture and sometimes of coaltar dyes. In these cheap extracts the per cent of alcohol often runs below 40, and as little as 4.5% by volume of alcohol has been found by the author in a commercial extract. With less than 45% of alcohol by volume, no appreciable amount of oil is dissolved, only a portion of citral, though such preparations are sometimes bottled as "pure extract of lemon." Time and again manufacturers have protested to the author that the purest oil was used by them, when notified that their brand contained no oil, or when prosecuted in court, and were with difficulty convinced that the trouble with their goods was that, on account of weak alcohol employed, the lemon oil used failed to get into the final product. It is true that a certain taste or odor of the lemon is present, even in cheap varieties wherein no oil is found, due to the fact that even dilute alcohol, when slowly percolating through the magnesia in which the oil is finely distributed, does abstract therefrom a certain amount of citral, which is, however, but a mere shadow of the substance and body possessed by a strong alcoholic solution of oil of lemon.

In many instances, where formulas appear stating the name and per cent of ingredients, these formulas are entirely deceptive and misleading, in that the statements are not borne out on analysis.

The flavor of the cheap extracts is sometimes reinforced by the addition of such substances as citral, oil of citronella, and oil of lemongrass, but minute quantities only of these pungent materials can be used, not exceeding 0.33% in the case of citral, and 0.1% in the case of the two last mentioned oils. Cane sugar and glycerin are sometimes found.

U. S. P. tincture of lemon peel owes its color to natural substances extracted by the alcohol. This color, however, readily fades on exposure to light. Other coloring matters employed are largely coal-tar dyes, and occasionally tincture of turmeric or saffron.

During 1901 practically all the brands of lemon extract sold in Massachusetts were collected and analyzed. 167 samples were examined, representing ahout 100 brands, and 139 samples were classed as adulterated, based on 5% lemon oil as a standard, and depending on whether or not the contents conformed to the labels on the bottles.

The typical analyses, given in tables on p. 864, are selected from the tabulated results of the above examination.*

Forty-two samples contained no lemon oil, ranging in content of alcohol from 4 to 45 per cent.

METHODS OF ANALYSIS OF LEMON EXTRACT.

A. S. Mitchell was the earliest among food chemists to systematically examine lemon extract, and to him are due the methods for determining oil of lemon, as well as various other tests now adopted provisionally by the A. O. A. C.†

Detection of Oil of Lemon.—If on adding a large excess of water to a little of the extract in a test-tube no cloudiness occurs, the oil may

^{*}An. Rep. Mass. State Board of Health, 1901, p. 459; Food and Drug Reprint, p. 41. † Jour. Am. Chem. Soc., 21, 1899, p. 1132; U. S. Dept. of Agric., Bur. of Chem., Bul. 65, p. 73; Bul. 107 (rev.), p. 159.

LEMON EXTRACTS OF STANDARD QUALITY.

Polarization	Lemon Oil,	Specific	Alcohol,	Foreign Ingredients.
in 200-mm.	Per Cent by	Gravity at	Per Cent by	
Tube.	Volume.	15.6° C.	Volume.	
30.8	9.1	0.8280	84.39	Turmeric
26.0	7.6	0.8402	80.49	Dinitrocresol
23.5	6.9	0.8352	81.74	
21.8	6.4	0.8396	82.88	
20.0	5·9	0.8335	84.24	
18.0	5·3	0.8268	• 86.82	
17.0	5.0	0.8496	80.06	

INFERIOR OR ADULTERATED LEMON EXTRACTS.

Polarization in 200-mm. Tube.	Lemon Oil, Per Cent by Volume.	Specific Gravity at 15.6° C.	Alcohol, Per Cent by Volume.	Foreign Ingredients.
14.0	4.1	0.8592	77.62	Dinitrocresol
12.2	3.6	0.8644	76.08	
11.0	3.1	0.8620	77.50	A coal-tar dye
9-9	2.9	0.8615	77.90	1
8. 6	2.3	0.8531	Šī.ģī	Dinitrocresol
6.8	2.0	0.8416	87.55	Tropæolin
5.0	1.5	0.8832	71.10	16
3.5	1.0	0.8939	67.68	1
2.8	0.8	0.8995	65.23	Dinitrocresol
2.2	0.6	0.8941	67.69	"
1.4	0.4	0.9136	59.40	A nitro dye
0.3	0.1	0.9408	46.40	Dinitrocresol
0.0	0.0	0.9937	4-49	Tropæolin
- 8.0	0.0	•••••		Invert sugar
27.0	0.0	• • • • • •	27-49	Cane sugar
0.0	0.0	•••••	47-35	Oil other than lemon

fairly be inferred to be absent. The degree of cloudiness produced is proportional to the amount of lemon oil present.

Determination of Lemon Oil.—Mitchell's Methods.—(1) By Polarization.—Polarize the undiluted extract in a 200-mm. tube at 20° C. Divide the reading on the Ventzke cane sugar scale by 3.4, and if cane sugar or other optically active substances are absent, the quotient expresses the per cent of lemon oil by volume. With instruments reading in circular degrees, divide the rotation in minutes at 20° C. by 62.5. If the Laurent instrument with sugar-scale is used, divide the sugar-scale reading by 4.8.

Cane sugar, though rarely found in lemon extract, is occasionally used in small amount. It is said to aid in the solution of the oil. If it is present, wash the solid residue from 10 cc. of the sample (dried on a water-bath) with three portions of 5 cc. each of ether, to remove waxy

and fatty matters, dry and weigh the residue of cane sugar, deducting 0.38 from the reading for each 0.1% of sugar so found.

(2) By Precipitation.—Transfer by a pipette 20 cc. of the extract to a Babcock milk-flask, add 1 cc. of dilute hydrochloric acid (1:1); add 25 to 28 cc. of water previously warmed to 60° C.; mix, and stand in water at 60° for five minutes; whirl in a centrifuge for five minutes; fill with warm water to bring the oil into the graduated neck of the flask, and repeat the whirling for two minutes; stand in water at 60° for a few minutes, and read the per cent of oil by volume. Where the oil of lemon is present in amounts over 2%, add to the percentage of oil found 0.4% to correct for the oil retained in solution. Where less than 2% and more than 1% is present, add 0.3% for correction.

Save the precipitated oil for the determination of refraction.

When the extract is made in accordance with the U. S. Pharmacopæia, the results by the two methods just given should agree within 0.2%.

To obtain per cent by weight from per cent by volume, as found by either of the above methods, multiply the volume percentage by 0.86, and divide the result by the specific gravity of the original extract.

Howard's Modification of Mitchell's Precipitation Method.*—Pipette 10 cc. of the extract in a Babcock milk bottle, and add in the following order, 25 cc. of cold water, 1 cc. hydrochloric acid (specific gravity 1.2), and 0.5 cc. chloroform. Close the mouth of the bottle with the thumb, and shake vigorously for not less than one minute. Whirl the bottle in a centrifuge for one and one-half to two minutes, thus forcing the chloroform and oil to the bottom of the bottle, and remove all but 3 or 4 cc. of the clear supernatant liquid by means of a glass tube of small bore connected with an aspirator.

To the residue add I cc. of ether, agitate thoroughly, plunge the bottle to the neck in a boiling-water bath, holding at slight angle, and rotate in the bath for exactly one minute. This step is best carried out by removing one of the small rings from a water- or steam-bath and holding the bottle in the live steam. The ether serves the purpose of steadily and rapidly sweeping out every trace of chloroform without appreciable loss of oil. Finally, cool the bottle, fill nearly to

^{*} Jour. Am. Chem. Soc., 30, 1908, p. 608.

the top of the neck with water at room temperature, centrifuge for one-half minute, read the column of separated oil to the top meniscus, and multiply the reading by two, thus obtaining the per cent of oil.

This method may also be used for determining the oil in extracts of orange, peppermint, clove, cinnamon, and cassia, employing in the case of the heavier oils dilute sulphuric acid (1:2), instead of water, in filling the bottles before the last centrifuging.

Determination of Alcohol.—Mitchell has shown that the difference in specific gravity between oil of lemon and stronger alcohol is not so great, but that a very close approximation to the true percentage of alcohol in lemon extracts may be obtained from the specific gravity itself, assuming, of course, that foreign substances, such as sugar, glycerin, etc., are absent. In the absence of such foreign substances determine the specific gravity of the sample, ascertain from the alcohol tables on pages 661–664, the per cent of alcohol by volume corresponding. This gross figure includes the lemon oil, the per cent of which should be deducted for the correct per cent of alcohol.

In the absence of oil of lemon, a measured portion of the original sample may be distilled, and the percentage of alcohol determined from the distillate in the usual manner, but when lemon oil is present, this should first be removed by diluting 50 cc. of the extract with water to 200 cc. exclusive of the oil in the sample, and shaking the mixture with 5 grams of magnesium carbonate in a flask, filtering through a dry filter, and determining the alcohol by distillation in a portion of the filtrate. The result is multiplied by 4 to correct for the dilution.

Determination of Citral.—Chace's Method.*—1. Reagents.—(a) Aldehyde-free Alcohol.—Allow alcohol (95% by volume) containing 5 grams of metaphenylene diamine hydrochloride per liter to stand for twenty-four hours with frequent shaking. Nothing is gained by previous treatment with potassium hydroxide. Boil under a reflux cooler for at least eight hours, longer if necessary, allow to stand over night and distil, rejecting the first 10 and the last 5 per cent which come over. Store in a dark, cool place in well-filled bottles. 25 cc. of this alcohol, on standing for twenty-minutes in the cooling bath with the fuchsin solution

^{*} Jour. Am. Chem. Soc., 28, 1906, p. 1472. U. S. Dept. of Agric., Bur. of Chem., Bul. 122, p. 32.

- (20 cc.), should develop only a faint pink coloration. If a stronger color is developed, treat again with metaphenylene diamine hydrochloride.
- (b) Fuchsin Solution.—Dissolve 0.5 gram of fuchsin in 250 cc. of water, add an aqueous solution of sulphur dioxide containing 16 grams of the gas, and allow to stand until colorless, then make up to 1 liter with distilled water. This solution should stand twelve hours before using, and should be discarded after three days.
- (c) Standard Citral Solution.—Use 1 mg. of c. p. citral per cc. in 50% by volume aldehyde-free alcohol. This solution deteriorates on standing, and should not be kept over three or four days.
- 2. Apparatus.—(a) A Cooling Bath.—Keep at from 14 to 16° C. The aldehyde-free alcohol, fuchsin solution, and comparison tubes are to be kept in this bath.
- (b) Colorimeter.—Any form of colorimeter, using a large volume of solution and adapted to rapid manipulation, may be used.

The comparison may also be made in Nessler or Hehner tubes.

3. Manipulation.—Weigh in a stoppered weighing flask approximately 25 grams of extract, transfer to a 50-cc. flask, and make up to the mark at room temperature with aldehyde-free alcohol. Measure at room temperature and transfer to a comparison tube 2 cc. of this solution. Add 25 cc. of the aldehyde-free alcohol (previously cooled in a bath), then 20 cc. of the fuchsin solution (also cooled), and finally make up to the 50-cc. mark with more aldehyde-free alcohol. Mix thoroughly, stopper, and place in the cooling bath for fifteen minutes. Prepare a standard for comparison at the same time and in the same manner, using 2 cc. of the standard citral solution. Remove and compare the colors developed. Calculate the amount of citral present and repeat the determination, using a quantity sufficient to give the sample approximately the strength of the standard. From this result calculate the amount of citral in the sample. If the comparisons are made in Nessler tubes, standards containing 1, 1.5, 2, 2.5, 3, 3.5, and 4 mg. should be prepared, and the trial comparison made against these, the final comparison being made with standards between 1.5 and 2.5 mg., varying but 0.25 mg.

It is absolutely essential to keep the reagents and comparison tubes at the required temperature. Comparisons should be made within one minute after removing the tubes from the bath. Where the comparisons are made in the bath (this being possible only where the bath is glass), the standards should be discarded within twenty-five minutes after

adding the fuchsin solution. Give samples and standards identical treatment.

Hiltner's Method.*—I. Reagents.—(a) Metaphenelene Diamine Hydrochloride Solution.—Prepare a 1% solution of metaphenelene diamine hydrochloride in 50% ethyl alcohol. Decolorize by shaking with fuller's earth or animal charcoal, and filter through a double filter. The solution should be bright and clear, free from suspended matter and practically colorless. It is well to prepare only enough solution for the day's work, as it darkens on standing. The color may be removed from old solutions by shaking again with fuller's earth.

- (b) Standard Citral Solution.—Dissolve 0.250 gram of c. p. citral in 50% ethyl alcohol and make up the solution to 250 cc.
- (c) Alcohol.—For the analysis of lemon extracts, 90 to 95 per cent alcohol should be used, but for terpeneless extracts alcohol of 40 to 50 per cent strength is sufficient. Filter to remove any suspended matter. The alcohol need not be purified from aldehyde. If not practically colorless, render slightly alkaline with sodium hydroxide and distil.
- 2. Apparatus.—The Schreiner colorimeter (page 77) or Eggertz tubes may be used. With this latter apparatus, alcohol is added, small quantities at a time, to the stronger colored solution until after shaking and viewing transversely, the colors in the two tubes are exactly matched. Calculations are then made by establishing a proportion between the volumes of samples taken and the final dilutions.
- 3. Manipulation.—All of the operations may be carried on at room temperature. Weigh into a 50-cc. graduated flask 25 grams of the extract, and make up to the mark with alcohol (90-95 per cent). Stopper the flask and mix the contents thoroughly. Pipette into the colorimeter tube 2 cc. of this solution, add 10 cc. of metaphenylene diamine hydrochloride reagent, and complete the volume to 50 cc. (or other standard volume) with alcohol. Compare at once the color with that of the standard, which should be prepared at the same time, using 2 cc. of standard citral solution and 10 cc. of the metaphenylene diamine reagent, and making up to standard volume with alcohol. From the result of this first determination, calculate the amount of standard citral solution that should be used in order to give approximately the same citral strength of the sample under examination, then repeat the determination.

^{*} A. O. A. C. Proc., 1908, U. S. Dept. of Agric., Bur. of Chem., Bul. 122, p. 34. Jour. Ind. Eng. Chem., I, 1909.

Methyl Alcohol has been used by unscrupulous manufacturers in lemon extracts. It is detected and determined by the refractometer method of Leach and Lythgoe (page 749).

As a confirmatory test for methyl alcohol the distillate, after testing by the Leach and Lythgoe method, may to advantage be subjected to the method of Mulliken and Scudder,* which depends on the conversion of the methyl alcohol to formaldehyde. The latter method is also useful as a rough preliminary test on the original extract without distillation, the extract, being, however, first diluted until the liquid contains approximately 12% by weight of alcohol, shaking with magnesium carbonate, and filtering when lemon oil is present.

Oxidize 10 cc. of the liquid in a test-tube as follows: Wind copper wire 1 mm. thick upon a rod or pencil 7 to 8 mm. thick, in such a manner as to inclose the fixed end of the wire, and to form a close coil 3 to 3.5 cm. long. Twist the two ends of the wire into a stem 20 cm. long, and bend the stem at right angles about 6 cm. from the free end, or so that the coil may be plunged to the bottom of a test-tube, preferably about 16 mm. wide and 16 cm. long. Heat the coil in the upper or oxidizing flame of a Bunsen burner to a red heat throughout. Plunge the heated coil to the bottom of the test-tube containing the diluted alcohol. Withdraw the coil after a second's time and dip it in water. Repeat the operation from three to five times, or until the film of copper oxide ceases to be reduced. Cool the liquid in the test-tube meanwhile by immersion in cold water.

Test for Formaldehyde.—Divide the oxidized liquid in the test-tube into two parts, testing one for formaldehyde with pure milk by the hydrochloric acid and ferric chloride test. Test the other portion by Mulliken and Scudder's resorcin test for formaldehyde, page 820, avoiding an excess of the reagent.†

Tests for Colors.—Evaporate a portion of the sample to dryness, dissolve the residue in water, and extract coal-tar colors if present by Arata's method, page 794, or with hydrochloric acid.

Much information may often be gained by treatment of the original extract with strong hydrochloric acid. If the color employed be turmeric, no change in color will be evident on addition of the acid. If tropæolin or methyl orange is present, the solution will turn pink, while partial decoloration of the solution indicates naphthol yellow S, and complete decoloration shows presence of dinitrocresols or naphthol yellow.

^{*} Am. Chem. Jour., 23, 1899, p. 266.

[†] Ibid, 24, 1900, p. 451.

Test for turmeric by boric acid, page 789.

Determination of Total Solids and Ash.—Total Solids are estimated by evaporating on the water-bath 10 grams of the sample in a tared dish, and drying at 100° to constant weight. If glycerin be present, it is difficult if not impossible to get a constant weight. Cane sugar and glycerin, if present, will be apparent in the residue. If capsicin has been used, it will be noticed by the taste.

Burn to an ash the residue from the solids in a muffle at a low red heat, cool in a desiccator, and weigh.

Glycerin is determined as in wine, page 703.

Detection of Tartaric or Citric Acid.—To a portion of the extract in a test-tube add an equal volume of water to precipitate the oil. Filter and add one or two drops of the filtrate to a test-tube half full of cold, clear lime water. If tartaric acid is present, a precipitate will come down, which is soluble in an excess of ammonium chloride or acetic acid.

Filter off the precipitate, or, if no precipitate is visible, heat the contents of the tube. Citric acid will precipitate in a large excess of hot lime water.

Examination of Lemon Oil.—The oil separated from the extract in the process of determining the lemon oil by precipitation (p. 865), is most readily examined for its purity, after drying with calcium chloride, by determination of its specific gravity, its index of refraction, or its refractometric reading with the Zeiss butyro-refractometer, and its polariscopic reading.

The specific gravity and refractometric readings are determined as with fixed oils, using with the butyro-refractometer a sodium flame or yellow bichromate color-screen, which gives perfectly sharp readings without dispersion.

The first table on page 871 shows readings on the Zeiss butyrorefractometer of pure lemon oil at various temperatures, using the sodium light.

For examination of high polarizing essential oils like oil of lemon, the author employs a 50-mm. tube, in order to get readings on the undiluted oil well within the limits of the cane sugar scale on the polariscope. If such a tube is not available, dilute the oil with an equal volume of alcohol, and use the 100-mm. tube. The second table on page 871 expresses constants of pure lemon oils and of various commonly employed adulterants, as determined in the laboratory of the Mascachusetts State Board of Health.

READINGS ON ZEISS BUTYRO-REFRACTOMETER OF LEMON OIL.

Tempera- ture, Centigrade.	Scale Reading.	Tempera- ture, Centigrade,	Scale Reading.	Tempera- ture, Centigrade.	Scale Reading.	Tempera- ture. Centigrade.	Scale Reading.
40.0	59-4	35.0	62.8	30.0	66.3	25.0	69.7
39-5	59-7	34-5	63.1	29.5	66.6	24.5	70.0
39.0	бо. 1	34.0	63.5	29.0	67.0	24.0	70.4
38.5	60.4	33-5	63.8	28.5	67.3	23.5	70.7
38.0	60.8	33.0	64.2	28.0	67.7	23.0	71.1
37-5	61.0	32-5	64.5	27.5	68.0	22.5	71.4
37.0	61.5	32.0	64.9	27.0	68.4	22.0	71.8
36.5	61.8	31.5	65.1	26.5	68.7	21.5	72.1
36.0	62.1	31.0	65.6	26.0	69.0	21.0	72-5
35-5	62.4	30.5	65.9	25.5	69.3	20.5	72.8
35.0	62.8	30.0	66.3	25.0	69.7	20.0	73.2

CONSTANTS OF SOME ESSENTIAL OILS.

. 01		fractometer Light) at—	Rotation in 100- Millimeter	Specific
Oil.	Temp.	Reading.	Tube, Ventzke Scale.	Gravity at 15.6° C.
Oil of lemon (lowest) """ (highest) """ grass (A. Giese) ""citronella (A. Giese) Terpeneless oil of lemon (Hansel's) """ grass (Hansel's) Citral (A. Giese)	25. 25. 22.5 22.5 23. 23. 22.5	69.5 71.2 96.9 87.1 87.9 91.0	173.0 184.5 -10.8 -10.2 -22.0 -5.6 -3.6	0.8580 0.8610 0.9309 0.9437 0.9463 0.9232 0.9296

Oil of Lemon is a light-yellow liquid, having the pleasant odor of fresh lemons, and an aromatic, mild, somewhat bitter after taste. It is obtained from the grated rind of the lemon either by treatment with hot water, skimming off the oil which rises to the surface, or by pressure, or by distillation with water. It is rapidly changed by action of air and light, becoming "terpeney," and under these conditions its solubility in alcohol seems to increase. Its composition is somewhat uncertain, but according to Wallach * nearly 90% consists of hydrocarbons, mostly terpenes, the most important of which is the terpene limonene † of the dextro-gyrate variety, also known as citrene.

Another important constituent of lemon oil is the aldehyde citral,

^{*} Liebig's Annalen, 227, p. 290.

[†] There are two limonenes, one of which is dextro- and the other lævo-rotary. The two are completely alike in their behavior, differing only in their optical rotation.

present to the extent of from 7 to 10 per cent. To this the odor of the oil is largely due. A second aldehyde, citronellal, is also present.

A frequent adulterant of lemon oil is turpentine oil, which lowers the rotation considerably, and is thus most easily rendered apparent.

Chace * detects small quantities of turpentine by the difference in crystalline form of pinene nitroso chloride from that of limonene nitroso chloride.

Citral (C₁₀H₁₆O) is an aldehyde present in lemon oil and in oil of lemon-grass, and, while it may be separated from these oils, is prepared artificially by oxidizing geraniol with chromic acid.† It is a mobile oil, and when perfectly pure is optically inactive. The commercial citral is, however, slightly lævo-rotary, due no doubt to impurities.

Oil of Lemon-grass is distilled from lemon-grass, Andropogon citratus (D. C.), cultivated in India. It is reddish yellow in color, and has an intense lemon-like odor and taste. Very little is known of its composition, but it seems to contain several aldehydes, one of which is citronellal, and another citral. The latter, however, is its chief constituent, being present to the extent of 70 to 75 per cent.

Citronellal $(C_{10}H_{18}O)$ is an aldehyde found in various oils, especially in citronella oil, from which it is readily separated. It is made artificially by the oxidation of the primary alcohol citronellol $(C_{10}H_{20}O)$. It is quite strongly dextro-rotary.

Oil of Citronella is distilled from the grass Andropogon nardus (L.), growing chiefly in Ceylon, India, and tropical East Africa. It is a yellowish-brown liquid with a pleasant and lasting odor. Citronellal is present in this oil to the extent of from 10 to 20 per cent, and the oil contains also from 10 to 15 per cent of terpenes, among which are camphene.

Tests for Citral, Citronellal, and Limonene.‡—Shake 2 cc. of the sample to be examined in a corked test-tube with 5 cc. of a solution of 10 grams of mercuric sulphate in sufficient 25% sulphuric acid to make 100 cc. Citral yields a bright-red color, which rapidly disappears, leaving a whitish compound, which floats on top. Citronellal forms a bright-yellow color, rem. ining for some time. Limonene forms an evanescent, faint flesh color, and leaves a white compound.

^{*} Jour. Am. Chem. Soc., 30, 1908, p. 1475.

[†] Tiemann, Berichte, 31, p. 3311.

[‡] Burgess, Chem. and Drugg., 57, p. 732.

ORANGE EXTRACT.

Orange Oil is a yellowish liquid, having the characteristic odor of orange, and a mild aromatic taste. It is prepared from orange peel in an analogous manner to that of lemon oil, which it somewhat resembles in chemical composition. At least 90% of orange oil, according to Walach, consists of dextro-limonene (citrene). It has a much higher specific rotatory power than lemon oil.

U. S. Standards.—Oil of Orange is the volatile oil obtained, by expression or alcoholic solution, from the fresh peel of the orange (Citrus aurantium L.) and has an optical rotation at 25° C. of not less than +95° in a 100-mm. tube.

Terpeneless Oil of Orange is oil of orange from which all or nearly all of the terpenes have been removed.

Orange Extract is the flavoring extract prepared from oil of orange, or from orange peel, or both, and contains not less than 5% by volume of oil of orange.

Terpeneless Extract of Orange is the flavoring extract prepared by shaking oil of orange with dilute alcohol, or by dissolving terpeneless oil of orange in dilute alcohol, and corresponds in flavoring strength to orange extract.

Analysis of Orange Extract.—The methods described under lemon extract (pp. 863 to 872) are also adapted for the analysis of orange extract.

In the determination of orange oil by Mitchell's polariscopic method divide the direct reading on the Ventzke scale, calculated for the 200-mm. tube, by 5.3 to obtain the per cent of orange oil by volume. To obtain the per cent by weight, multiply the per cent by volume by 0.85 and divide by the specific gravity of the extract.

ALMOND EXTRACT.

Oil of Bitter Almonds is obtained by distilling crushed bitter almonds, peach seeds, or apricot seeds with water. It should be remembered that both sweet and bitter almonds yield a bland fixed oil on pressure, which is not to be confounded with the volatile oil yielded on distillation of the bitter almonds after the fixed oil has been pressed out. Bitter almonds contain a glucoside, amygdalin, together with a ferment known as emulsin or synaptase, which, acting on the amygdalin in the distillation, produces benzaldehyde and hydrocyanic acid as follows:

The unpurified oil of bitter almonds consists largely of benzaldehyde, with a small amount of the poisonous hydrocyanic acid. Nearly all of the commercial oil is made from the cheaper apricot and peach seeds rather than those of the bitter almond, but the product is practically identical. The oil is freed from hydrocyanic acid by agitating with calcium hydrate and a solution of ferrous chloride, distilling the mixture, and drying the oil which comes over with calcium chloride.

Benzaldehyde constitutes 90 to 95 per cent of oil of bitter almonds, having a bitter, acrid, burning taste, and a marked almond odor. The specific gravity of the crude oil varies from 1.052 to 1.082, while that of the purified oil (benzaldehyde) at 20° is 1.0455. Its boiling-point is 180° C. On standing it becomes readily oxidizable to benzoic acid. It is readily soluble in alcohol and ether. Its solubility in water is slight, 1:300. Its index of refraction at 20° C. is 1.5446. It should be noted that the refractive indices of almond oil, whether with or without hydrocyanic acid, and of artificial benzaldehyde are nearly the same.

Benzaldehyde is produced artificially in a variety of ways, but is chiefly prepared by the action of chlorine on hot toluene. The resulting benzyl chloride is distilled with lead nitrate and water in an atmosphere of carbon dioxide, which forms benzoic aldehyde. Synthetic benzaldehyde has the same properties as the purified oil of bitter almonds, and has largely displaced it in the market, not the least of its advantages being its freedom from hydrocyanic acid.

Almond Extract.—Essence of bitter almonds, or Spiritus amygdalæ amaræ, is thus prepared according to the U. S. Pharmacopæia:

Oil of bitter almonds	IO CC.
Alcohol	800 cc.
Distilled water sufficient to make	1000 CC.

Thus 1% of almond oil is present in the product.

U. S. Standards.—Oil of Bitter Almonds, commercial, is the volatile oil obtained from the seed of the bitter almond (Amygdalus communis L.), the apricot (Prunus armeniaca L.), or the peach (Amygdalus persica L.).

Almond Extract is the flavoring extract prepared from oil of bitter

almonds, free from hydrocyanic acid, and contains not less than 1% by volume of oil of bitter almonds.

Adulteration of Almond Oil.—The official essence of the Pharma-copœia does not specify that the almond oil used be perfectly free from hydrocyanic acid, in spite of the fact that its highly poisonous nature is well known, and that it exists in the crude oil to the extent of from 4 to 6 per cent. True, but little of it is found in the extract, but in these days, when the unannounced presence in foods of such substances as antiseptics and coloring matters is regarded as questionable from a sanitary standpoint, in spite of the fact that their toxic effects on man are still matters of controversy, there thould be little hesitancy in pronouncing the presence of prussic acid objectionable, especially when a pure almond oil entirely free from it is readily obtainable.

The presence of nitrobenzol or oil of mirbane as a substitute of almond oil is to be looked for. This substance is sometimes, though incorrectly, called artificial oil of bitter almonds. It is a heavy, yellow liquid of the composition C₆H₅NO₂, readily soluble in water. Its specific gravity at 20° C. is 1.2039. Its boiling-point is 205° C. It is formed by the action of nitric acid on benzol. It possesses a highly pungent odor, somewhat like that of oil of bitter almonds, though more penetrating and less refined. Its index of refraction at 20° C. is 1.5517.

METHODS OF ANALYSIS OF ALMOND EXTRACT.

Determination of Benzaldehyde.—Hortvet and West's Method.*—Measure 10 cc. of the extract into a 100-cc. flask, add 10 cc. of a 10% sodium hydroxide solution, and 20 cc. of a 3% hydrogen peroxide solution, cover with a watch-glass and place on a water-oven. Oxidation begins almost immediately and should be continued from five to ten minutes after all odor of benzaldehyde has disappeared, which usually requires from twenty to thirty minutes. If nitrobenzol be present, it will be indicated at this point by its odor. When the oxidation of the aldehyde is complete, remove the flask from the water-oven, transfer the contents to a separatory funnel, rinsing off the watch-glass, add 10 cc. of a 20% sulphuric acid solution, and cool the contents of the funnel to room temperature under the water tap. Extract the benzoic acid with three portions of 50, 30, and 20 cc. of ether, respectively, wash the combined extracts in another separatory funnel with two portions of from 25 to

^{*} Jour. Ind. Eng. Chem., 1, 1909, p. 86.

30 cc. of distilled water, or until all the sulphuric acid is removed. Filter into a tared dish, wash with ether, allow to evaporate at room temperature, and finally dry over night in a desiccator, and weigh. The per cent of benzaldehyde (B) is obtained from the weight of the acid (W) by the following formula:

$$B = \frac{0.869 \times 10 \times W}{1.045}.$$

If desired the benzoic acid may be titrated, and the benzaldehyde calculated from the amount of standard alkali required for neutralization. The process is as follows: Dissolve the benzoic acid obtained as above described, except that it need not be dried in a desiccator, in 95% alcohol made neutral to phenolphthalein with tenth-normal sodium hydroxide, dilute with an equal volume of water, and titrate with tenth-normal sodium hydroxide, using phenolphthalein as indicator. The per cent of benzaldehyde (B) is calculated from the cc. of tenth-normal alkali (V) by the following formula:

$$B = \frac{V \times 0.01061 \times 10}{1.045}.$$

Detection of Nitrobenzol.*—Boil 15 cc. of the extract in a test-tube with a few drops of a strong solution of potassium hydroxide. Nitrobenzol produces a blood-red coloration.

Distinction between Benzaldehyde and Nitrobenzol.—Treat 20 cc. of the extract with 5 to 10 cc. of a cold, saturated aqueous solution of sodium bisulphite in a test-tube, and shake vigorously. Transfer to an evaporating-dish, and heat on the water-bath till the alcohol is driven off. At this stage benzaldehyde remains in the hot solution as a crystal-line salt, and the solution gives off no almond odor.

Nitrobenzol, on the contrary, does not combine with the bisulphite and is insoluble, forming globules of oil on the surface of the hot liquid, and in addition giving off the pungent odor so characteristic of the substance.

Separation of Nitrobenzol and Benzaldehyde.—If by the qualitative test nitrobenzol is found, shake vigorously as before 50 cc. of the extract with 10 cc. of the saturated sodium bisulphite solution in a corked flask, and transfer with 100 cc. of water to a large separatory funnel. Shake out the nitrobenzol from the solution with four successive portions of

^{*} Holde, Jour. Soc. Chem. Ind., 13, 1893, p. 906.

petroleum ether of 15 to 20 cc. each, and after washing with water the combined petroleum ether, transfer it to a tared dish, in which it is allowed to evaporate spontaneously.

It is extremely difficult to avoid loss of some of the nitrobenzol by this process, but even if the weighed residue fails to show the full amount originally used, enough will usually be extracted to admit of testing on the refractometer, and of otherwise verifying its character.

After removal of the nitrobenzol, make the residual solution in the separatory funnel strongly alkaline with sodium hydroxide, and shake out the benzaldehyde, if present, with petroleum ether as previously described. If after making the solution alkaline no odor of benzaldehyde is apparent, the absence of benzaldehyde may be inferred.

Distinction between Artificial Benzaldehyde and Pure Almond Oil.— Test the final residue from the ether extract by shaking with an equal volume of concentrated sulphuric acid in a test-tube. With natural oil of almonds a clear, brilliant, but dark currant-red color is produced, while with artificial benzaldehyde, the acid produces a dirty brown color with the formation of a precipitate.

Determination of Alcohol.—In the absence of other flavoring substances than nitrobenzol and benzaldehyde, which are rarely present to an extent exceeding 1%, a sufficiently close approximation for most purposes can be gained by estimating the alcohol from the direct specific gravity of the extract.

Detection of Hydrocyanic Acid.—To a few cubic centimeters of extract in a test-tube add a few drops of a mixture of solutions of ferrous sulphate and ferric chloride, the ferrous salt being in excess. Make alkaline with sodium hydroxide, and add enough dilute hydrochloric acid to dissolve the precipitate formed by the alkali. Presence of a blue coloration or precipitate, due to the formation of Prussian blue, indicates hydrocyanic acid. The reaction is very delicate.

Determination of Hydrocyanic Acid.*—Hydrocyanic acid may be determined by titration with tenth-normal silver nitrate solution. 25 cc. of the extract are measured into a flask, and 5 cc. of freshly prepared magnesium hydroxide suspended in water are added, or enough to make the reaction alkaline.

A few drops of a solution of potassium chromate are then introduced, and the tenth-normal silver nitrate solution added till, with shaking, the formation of the red silver chromate indicates the end-point. I cc. of silver solution equals 0.0027 gram of hydrocyanic acid.

^{*} Vielhaber, Arch. Pharm. (3), 13, 408.

WINTERGREEN EXTRACT.

Wintergreen Oil.—True oil of wintergreen is obtained by distillation from the leaves of the wintergreen plant (Gaultheria procumbers L.). Gildermeister and Hoffman * state that the specific gravity at 15° is 1.180 to 1.187, the boiling-point 218 to 221° C. It is slightly leworotatory $(a_D = -0.0^{\circ} 25' \text{ to } -1^{\circ})$.

Oil of betula or sweet birch is distilled from the bark of the black birch (*Betula lenta L.*). It has the same specific gravity and boiling-point as oil of wintergreen, but unlike the latter is optically inactive. It differs somewhat from oil of wintergreen in taste and odor, but is hardly distinguishable in these respects from synthetic methyl salicylate.

Both oil of wintergreen and oil of sweet birch consist almost entirely of methyl salicylate, the former containing, according to Power and Kleber,† as much as 99.8% of this substance.

U. S. Standards.—Oil of Wintergreen is the volatile oil distilled from the leaves of the Gaultheria procumbers L.

Wintergreen Extract is the flavoring extract prepared from oil of wintergreen, and contains not less than 3% by volume of oil of wintergreen.

Spirit of Gautheria of the U. S. P. is a mixture of 50 cc. of oil of wintergreen and 950 cc. of alcohol. It accordingly contains 5% by volume of the essential oil.

Adulteration of Wintergreen Extract.—Synthetic methyl salicylate is very commonly substituted for both wintergreen and sweet birch oil, and sweet birch oil in turn for wintergreen oil. The production of true wintergreen oil is small, the so-called natural wintergreen oil of commerce being usually sweet birch oil. The sense of smell is the best means of distinguishing the two oils; polarization is of rather uncertain value, owing to low rotatory power of the wintergreen oil.

Determination of Wintergreen Oil.—Hortvet and West's Method.‡—Measure 10 cc. of the extract into a 100-cc. beaker, add 10 cc. of 10% potassium hydroxide solution, and heat the mixture over a boiling waterbath until the odor of oil of wintergreen has disappeared and the liquid is reduced to about one-half its original volume. By this treatment the methyl salicylate is converted into the potassium salt. Liberate the

^{*} The Volatile Oils. Translated by Kremers, Milwaukee, 1900, p. 588.

[†] Pharm. Rund., 13, p. 228.

[‡] Jour. Ind. Eng. Chem., 1, 1909, p. 90.

salicylic acid by the addition of an excess of 10% hydrochloric acid, cool, and extract in a separatory funnel with three portions of 40, 30, and 20 cc. of ether respectively. Pour the combined ether extracts through a dry filter into a weighed dish, wash the filter with 10 cc. of ether, evaporate filtrate and washings slowly at 50° C., dry one hour in a desiccator, and weigh. The per cent of wintergreen oil by volume (M) is obtained from the weight of salicylic acid (S) by the following formula:

$$M = \frac{1.101 \times 10 \times S}{1.18}.$$

Howard's Method.—Proceed as described on page 865, except that the heavy oil is brought into the graduated portion of the Babcock bottle by addition of dilute sulphuric acid (1:2), taking care that the acid is not over 25° C. and avoiding agitation.

PEPPERMINT EXTRACT

Peppermint Oil is obtained from various plants of the genus Mentha, which are commonly classed as sub-species or varieties of M. piperita. Owing in large part to the botanical differences in the plants from which it is made, peppermint oil from different regions differs greatly in its chemical and physical constants as shown by the following table compiled from figures given by Gildermeister and Hoffmann:*

	Specific Gravity.	Rotation, a_D .	Total Menthol, Per Cent.
American English Japanese Saxon German French Russian	0.900 to 0.910 0.895 to 0.900 0.900 to 0.915 0.899 to 0.930 0.918 to 0.920	-18° to -33° -22° to -33° -30° to -42° -25° to -33° -27° to -33° -5° to -9° -17° to -22°	48 to 60 56 to 66 70 to 91 54 to 68 43 to 46 50,2

U. S. Standards.—Peppermint is the leaves and flowering tops of Mentha piperita L.

Oil of Peppermint is the volatile oil obtained from peppermint, and contains not less than 50% by weight of menthol.

Peppermint Extract is the flavoring extract prepared from oil of pepper-

^{*} The Volatile Oils. Translated by Edward Kremers, Milwaukee, 1900.

mint, or from peppermint, or both, and contains not less than 3% by volume of oil of peppermint.

Analysis of Peppermint Extract.—Owing to the wide variation in the rotatory power of peppermint oil, only a roughly approximate idea of the oil content of peppermint extract can be gained by polarization. The variation in the percentage of menthol in the oil is also too great to permit of a method based on the amount of this constituent. Mitchell's precipitation method, as originally described (page 864), does not effect a complete separation of the oil, but Howard's modification (page 865) gives satisfactory results, and is well adapted for purposes of inspection.

SPEARMINT EXTRACT.

U. S. Standards.—Spearmint is the leaves and flowering tops of Mentha spicata L.

Oil of Spearmint is the volatile oil obtained from spearmint.

Spearmint Extract is the flavoring extract prepared from oil of spearmint, or from spearmint, or both, and contains not less than 3% by volume of oil of spearmint.

SPICE EXTRACTS.

Alcoholic solutions of the essential oils of spices are used to some extent instead of the spices themselves. The following are the definitions of these extracts and the oils from which they are prepared, as adopted by the joint committee on standards and the U. S. Secretary of Agriculture:

U. S. Standards.—Anise Extract is the flavoring extract prepared from oil of anise, and contains not less than 3% by volume of oil of anise.

Oil of Anise is the volatile oil obtained from the anise seed.

Celery Seed Extract is the flavoring extract prepared from celery seed or the oil of celery seed, or both, and contains not less than 0.3% by volume of oil of celery seed.

Oil of Celery Seed is the volatile oil obtained from celery seed.

Cassia Extract is the flavoring extract prepared from oil of cassia, and contains not less than 2% by volume of oil of cassia.

Oil of Cassia is the lead-free volatile oil obtained from the leaves or bark of Cinnamomum cassia Bl., and contains not less than 75% by weight of cinnamic aldehyde.

Cinnamon Extract is the flavoring extract prepared from oil of cinnamon, and contains not less than 2% by volume of oil of cinnamon.

Oil of Cinnamon is the lead-free volatile oil obtained from the bark of the Ceylon cinnamon (Cinnamonum zeylanicum Breyne), and contains not less than 65% by weight of cinnamic aldehyde and not more than 10% by weight of eugenol.

Clove Extract is the flavoring extract prepared from oil of cloves, and contains not less than 2% by volume of oil of cloves.

Oil of Cloves is the lead-free, volatile oil obtained from cloves.

Ginger Extract is the flavoring extract prepared from ginger, and contains in each 100 cc. the alcohol-soluble matters from not less than 20 grams of ginger.

Nutmeg Extract is the flavoring extract prepared from oil of nutmeg, and contains not less than 2% by volume of oil of nutmeg.

Oil of Nutmeg is the volatile oil obtained from nutmegs.

Savory Extract is the flavoring extract prepared from oil of savory, or from savory, or both, and contains not less than 0.35% by volume of oil of savory.

Oil of Savory is the volatile oil obtained from savory.

Star Anise Extract is the flavoring extract prepared from oil of star anise, and contains not less than 3% by volume of oil of star anise.

Oil of Star Anise is the volatile oil distilled from the fruit of the star anise (Illicium verum Hook).

Sweet Basil Extract is the flavoring extract prepared from oil of sweet basil, or from sweet basil, or both, and contains not less than 0.1% by volume of oil of sweet basil.

Sweet Basil, Basil, is the leaves and tops of Ocymum basilicum L.

Oil of Sweet Basil is the volatile oil obtained from basil.

Sweet Majoram Extract, Majoram Extract, is the flavoring extract prepared from the oil of majoram, or from majoram, or both, and contains not less than 1% by volume of oil of majoram.

Oil of Majoram is the volatile oil obtained from majoram.

Thyme Extract is the flavoring extract prepared from oil of thyme, or from thyme, or both, and contains not less than 0.2% by volume of oil of thyme.

Oil of Thyme is the volatile oil obtained from thyme.

Determination of Essential Oil in Cinnamon, Cassia, and Clove Extracts.—Howard's Method.—Proceed as with wintergreen extract, page 879.

Hortvet and West's Method.*—Place 10 cc. of the extract and 50 cc. of water in a separatory funnel, and extract with three portions of ether measuring respectively 50, 30, and 20 cc. Wash the combined extracts successively with 25 and 30 cc. of distilled water, and filter through a dry funnel into a wide-mouth flask, washing out the funnel and filter with a little ether. In the case of cinnamon extract, transfer the ether extract before filtering to a 150-cc. flask, shake for a few minutes with some granulated calcium chloride, then filter in the manner described. Evaporate off the ether as rapidly as possible on a boiling water-bath until only a few drops remain. At this point remove the flask from the bath, and rotate rapidly for a few minutes, spreading the residue over the sides of the flask. The rapid evaporation of the remaining ether cools the flask to near room temperature. When the odor of ether has disappeared, stopper the flask and weigh.

In the case of cassia and clove oils, where the ether extract is not first dried with calcium chloride, a slight cloudiness gathers on the flask as the last traces of ether disappear, due to the presence of a little moisture. In such case allow the flask to stand on the balance-pan until the film disappears, requiring not longer than two to three minutes, then stopper, and weigh.

The per cent of oil by volume (V) is calculated from the weight of oil (W) by the following formula:

$$V = \frac{100 \times W}{10 \times 1.050}.$$

The oil thus extracted may be used for determination of the refractive index. After dissolving in a little alcohol it may be tested with ferric chloride solution. By this test cinnamon oil gives a green, cassia oil a brown, and clove oil a deep blue, coloration.

Determination of Essential Oil in Nutmeg Extract.—Follow Mitchell's precipitation method, page 865.

ROSE EXTRACT.

U. S. Standards.—Rose Extract is the flavoring extract prepared from otto of roses, with or without red rose petals, and contains not less than 0.4% by volume of otto of roses.

Otto of Roses is the volatile oil obtained from the petals of Rosa damascena Mill., R. centifolia L., or R. moschata L.

^{*} Jour. Ind. Eng. Chem., 1, 1909, p. 88.

Determination of Rose Oil.—Hortvet and West's Method.*—Measure 25 cc. of the extract into a separatory funnel, add 50 cc. of water, mix thoroughly, acidify with 1 cc. of hydrochloric acid (1:1), and extract with three portions of 20 cc. each of ether. Transfer the combined ether extracts to a 150-cc. flask, shake for a few minutes with some granulated calcium chloride, allow to settle until clear, then decant through a dry filter into a flat bottom glass dish previously weighed together with a cover-glass. Wash the calcium chloride and filter twice with 10 cc. of ether, and add the washings to the glass dish. Cover the dish, place in a vacuum desiccator over sulphuric acid, allow to remain until all traces of ether and alcohol are removed, and weigh. Repeat the drying in the desiccator, for one hour periods, until the weight is practically constant. The final weight, divided by 0.86 and multiplied by 5, gives the per cent of oil of rose by volume.

IMITATION FRUIT FLAVORS.

Nearly all the fruits possess distinctive flavors, which are desirable in food preparations, and which may be made to impart their flavor to such substances as confections, ice cream, dessert mixtures, jellies, etc., by simply mixing with these foods the fresh or preserved fruit or fruit juice in sufficient quantity. In many cases, however, it is not found possible or practicable to prepare from the fruits themselves an extract sufficiently concentrated to give the distinctive fruit flavor, when used in moderate quantity, and hence the use of artificial fruit essences made up of compound ethers, mixed in varying combinations and proportions to imitate more or less closely various fruit flavors.

These ethers are usually much more pungent and penetrating than the fruits which they imitate, and, while lacking the delicacy and refinement of the original fruits, serve to impart a certain semblance of the genuine flavor in a convenient and highly concentrated form.

Some of the single compound ethers possess a remarkable resemblance to particular fruits, while to imitate other fruits a mixture of various ethers and flavoring materials, such as lemon and other volatile oils, vanilla, organic acids, chloroform, etc., is necessary. These artificial essences should in all cases be sold as such, and not as "pure fruit flavors."

^{*} Jour. Ind. Eng. Chem., 1, 1909, p. 89.

Imitation Pineapple Essence is made up by dissolving in alcohol butyric ether, $C_4H_7(C_2H_6)O_2$, which possesses a distinct pineapple flavor, and is prepared by mixing 100 parts of butyric acid $(C_4H_6O_2)$, 100 parts of alcohol, and 50 parts of sulphuric acid, and shaking. Butyric ether is sparingly soluble in water, and boils at 121° C.

Imitation Quince Essence depends as a basis on ethyl pelargonate, sometimes called pelargonic or cenanthic ether, C_2H_6 , $C_9H_{17}O_2$, dissolved in alcohol. Pelargonic ether is formed by digestion with the aid of heat of pelargonic acid and alcohol. Pelargonic acid, $C_9H_{18}O_2$, is first obtained by the action of nitric acid on oil of rue. Pelargonic ether is a colorless liquid, having a specific gravity of 0.8635 at 17.5° C. Its boiling-point is 227° to 228° C. It is insoluble in water.

Imitation Jargonelle Pear Essence consists of an alcoholic solution of amyl or pentyl acetate, C_5H_{11} , $C_2H_3O_2$. This is prepared by distilling a mixture of one part of amyl alcohol, two parts of potassium acetate, and one part of concentrated sulphuric acid. It is a colorless liquid, insoluble in water, and having a boiling-point of 137° C.

Imitation Banana Essence is made up of a mixture of amyl acetate and butyric ether.

Imitation Apple Essence is composed of an alcoholic solution of amyl valerianate, sometimes called apple oil, C_5H_{11} , $C_5H_9O_2$, prepared by mixing four parts of amyl alcohol with four of sulphuric acid, and adding the mixture when cold to five parts of valerianic acid. The specific gravity of amyl valerianate is 0.879 at 0° C. and its boiling-point is 188° C.

The table on p. 885, prepared by Kletzinsky, shows the composition of a large variety of these artificial essences. The numerals in the various columns indicate the parts by volume to be added to one hundred parts of deodorized alcohol.

Determination of Esters.—Add to 25 grams of the extract 2 cc. of sodium hydroxide solution (100 grams in 100 cc. of water), 100 cc. of water and heat under a reflux condenser one half-hour. Acidify with 5 cc. of dilute sulphuric acid (1:4), add a few pieces of pumice stone, distil in a current of steam and titrate the distillate with tenth-normal alkali, using phenophthalein as indicator. The number of cc. required represents the total volatile acids free and combined. Determine free volatile acids, if present by direct distillation and titration of the distillate. The difference between the two titrations is calculated as ethyl acetate.

COMPOSITION OF IMITATION ESSENCES.

	Chloroform.	Nitrous Ether.	Aldehyde.	Acetic Ether.	Formic Ether.	Butyric Ether.	Valerianic Ether.	Benzoic Ether.	Gnanthylic Ether.	Oil of Persi- cot.	Sebacic Ether.	Methylr salicylic Ether.
Pineapple Melon Strawberry Raspberry Gooseberry Grape. Apple Orange. Pear Lemon. Black cherry Cherry Plum Apricot. Peach Currant	1 2 1 2 1 1	1 I	1 1 1 2 2 2 2 2 5 2 1	5 5 5 10 10 5 5 5	1 1 1 1 1	5 4 5 1 1	5	1 1 5 5	1 10	2 4 5	10	I I
					υ							-
		Alcohol.	cetic	outyric	raleriani	Lemon.	Orange.		l	d Alcho	. 	d
		Amyi Alcohol.	Amyl-acetic Ether.	Amyl-butyric Ether.	Amyl-valerianic Ether.	Oil of Lemon.	Oil of Orange.	Tartaric Acid.	Acid.	Succinic Acid.	Benzoic oii	Glycerin.

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CHAPTER XXL

CANNED AND BOTTLED VEGETABLES, RELISHES, AND FRUIT PRODUCTS.

CANNED VEGETABLES AND FRUITS.

STRICTLY speaking all varieties of canned foods found in the market, whether meats, fruits, or vegetables, in order to be entirely beyond criticism, should not differ from the corresponding freshly cooked varieties which they are intended to replace, excepting that they are free from bacteria. Such a degree of perfection is, however, difficult, even if possible, to attain, and nearly all commercial canned products, even if made from the best materials, are liable to contain either antiseptic substances or coloring-matter intentionally added by the manufacturer, or metallic impurities accidentally derived from the vessels in which they are prepared, or from the containers in which they are sealed. In spite of these objections, canned foods form a convenient, and in some cases indispensable means of furnishing both necessities and luxuries for the table. The canning of foods is especially useful for preserving them during long periods of time, for enabling certain fruits and vegetables to be enjoyed out of season, and for furnishing supplies in a convenient manner to inaccessible places where fresh foods are not readily obtainable, as in the case of armies in the field, of vessels at sea, of campers in the woods, etc. Canned goods in great variety are used in nearly every household.

When it is considered that in the United States alone something like one hundred million cans of corn are packed in a single year, about the same quantity of peas, and one hundred and fifty million cans of tomatoes, to say nothing of an ever-increasing variety of other foods, some idea may be gained of the enormous proportions to which the canning industry has grown. It is comforting to know that, in view of their wide-spread consumption, the greater portion of such foods found on the market are

comparatively harmless, as is evidenced by the fact that few cases of injury to health have been directly traceable to their use.

Method of Canning Food.—Various modifications as to details exist with different products and in different localities, but in general the principle of canning in tin is the same in all cases. The fresh product is cleaned carefully, and packed in cans with the requisite amount of water. The cans are then sealed, and subjected to the effect of steam or boiling water till the contents are thoroughly cooked. Each can is then tapped or punctured at one end to expel the air, and again heated, after which the hole is closed by a lump of solder, thus forming a vacuum in the can, which is afterwards heated for a sufficient time to destroy the bacteria, usually for several hours.

The above mode of procedure is the time-honored one, and is still in vogue in most localities, but a more modern method, much in use at present, consists in first cooking the food at a temperature of 82° to 88° C. before transferring to the cans, and afterwards subjecting the cans when sealed to a high heat of about 125° C. in dry air in so-called retorts, this heating or "processing," as it is termed, being carried on for a sufficient length of time to completely sterilize the contents of the can. Obviously a much shorter time is required for this than when the temperature of boiling water is employed, and the sterilization is much more effective.

Cooked vegetables and fruit products put up in glass jars or bottles are tightly sealed when hot, either with screw-caps or with some form of cover held by a clamp, or with metal or hard-rubber caps fitting over a flanged mouth. Frequently a soft-rubber ring is inserted between the cover and the mouth of the jar or bottle. The material of the cover is generally either glass, porcelain, or metal. Cork stoppers are, however, sometimes pressed into the mouths of the bottles, and made extra tight therein with sealing-wax. These stoppers are occasionally soaked in paraffin. Thus the contents of the jar may be exposed to porcelain, glass, metal, rubber, or cork, according to the material of the cover and the method of sealing.

The preservation of food by canning was long thought to be due to the perfect exclusion of air, but is now known to depend on the perfect sterilization, or destruction of bacteria, and it has been proved that as far as keeping qualities are concerned, it makes no difference whether or not air is present in the can, if the contents are sterile, though for purposes of inspection the vacuum, in the case of tin cans, is of great use, in that as a natural consequence of the vacuum, when the goods are sound, the ends of the can are usually concave. The highest aim of the canner should be to retain in his product as far as possible the appearance, palatability, and nutritive value of the freshly cooked food.

DDOVIMATE	COMPOSITION	OF CANINED	VECETABLES	ABITA TODITIONS
PRUMMAIL	COMPOSITION	OF CANNED	VEGETABLES	AND FRUITS.

	No. of Analyses.	Water.	Protein.	Fat.	Total Carbohy- drates.	Crude Piber.	Ash.	Fuel Value per Pound.
CANNED VEGETABLES. Artichokes. Asparagus. Beans, baked. '' string. '' Lima. Brussels sprouts. Corn, green. Peas, green. Pumpkin. Squash. Succotash Tomatoes	3 14 21 29 16 1 52 88 7 5 12	92-5 94-4 68-9 93-7 79-5 93-7 76-1 85-3 91-6 87-6 75-9 94-0	.8 1.5 6.9 1.1 4.0 1.5 2.8 3.6 .9 3.6	.1 2.5 .1 .3 .1 1.2 .2 .2	5.0 2.8 19.6 3.8 14.6 3.4 19.0 9.8 6.7 10.5 18.6 4.0	.6 -5 2-5 -5 1.2 -8 1.2 1.1	1.7 1.2 2.1 1.3 1.6 1.3 .9 1.1	95 360 95 455 255 150 235 455 105
CANNED FRUITS. Apples, crab. Apple sauce. Apricots. Blackberries. Blueberries. Cherries Peaches. Pears. Pineapples. Strawberries.	1 1 3 1 3 4 1	42.4 61.1 81.4 40.0 85.6 77.2 88.1 81.1 61.8 74.8	.3 .2 .9 .8 .6 I.I .7 .3 .4	2.4 .8 2.1 .6 .1 .1 .3 .7	54.4 37.2 17.3 56.4 12.8 21.1 10.8 18.0 36.4 24.0		•5 •7 •4 •7 •4 •5 •3 •7 •5	1,120 730 340 1,150 275 415 220 355 715 460

^{*}U. S. Dept. of Agric., Exp. Sta. Bul. 28, p. 70.

Methods of Proximate Analysis.—As a rule, the contents of canned goods are intended to be entirely edible throughout, and contain little or no refuse or portions to be rejected. An exception to this is the occasional canning of certain fruits with stones or pits, which are, of course, to be removed. The can or package is first weighed before opening, and later the cleaned receptacle is weighed after its contents have been removed. The weight of the contents is thus ascertained by difference.

For the analytical determinations, the contents of the can or bottle are intimately mixed to form a homogeneous pulp, so that parts taken for analysis are fairly representative of the whole. If considerable liquid is present, with some solid masses held in suspension therein, the liquid is best drained off, and the solid portions pulped separately in any convenient manner, as by the use of a mortar, or by means of a household

food-chopper. The whole is then thoroughly mingled together. If desired, the weight of the liquid and solid portions may be separately ascertained before mixing.

The analyst should use judgment and discrimination as to how various portions of the mass are to be best measured out for the determinations. Much depends on the consistency of the pulpy mass. It is often convenient to make a 20% solution or mixture of the material with water, using say 50 grams of the pulped sample in 250 cc. of water, such of the sample as is insoluble being disintegrated by shaking.

Methods for determining water, ether extract, crude fiber, protein, and ash do not differ materially from those employed in the case of corresponding fresh fruits and vegetables.

These determinations, in the case of canned products, while useful in showing their food value, give little information as to their adulteration by the substitution of foreign vegetable or fruit pulp.

ACCIDENTAL IMPURITIES.—Under this head are included (1) products of decomposition, due to the incomplete sterilization of the contents of the can, and (2) metallic salts due to the solvent action of the juices of the contents on the inner surface of the can, or of the vessel in which the product has been previously cooked.

Decomposition, and the Detection of Spoiled Cans.—In the case of canned vegetable products, decomposition rarely results in the formation of ptomaines even after the can has long been open, though these toxins are sometimes formed, in canned meat and fish. Decomposition is readily apparent after opening a can, from a cursory examination of its contents. The appearance, taste, and odor will not fail to indicate the unfitness of the contents for food, if decomposition is at all advanced. It is, however, often of great advantage to detect spoiled cans without opening. As a rule, when a can is spoiled, it is usually in the condition termed "blown," i.e., with its ends convex, instead of normal or concave.

According to Prescott and Underwood,* although nearly all forms of bacterial decomposition are accompanied by bulging of the ends of the cans, there are some exceptions. In the souring of canned sweet corn,† for instance, it is exceptional that swelling occurs. Ordinarily, in the factory inspection of canned goods before shipping, not only are the bulged cans or "swells," as they are termed, sifted out, but the condition

^{*} Tech. Quart., 11, 1898, pp. 6-30; also 10, 1897, p. 183.

[†] These experimenters found at least twelve varieties of bacteria to which the souring of corn is apparently due.

of the cans is tested by sounding or striking the cans. If the contents are sweet, a peculiar note is produced when the can is struck, readily distinguishable from the dull tone of the unsound can by any one familiar with the work.

As stated above, concavity in the ends of the can indicates that the contents are in good condition.

Prescott and Underwood further state that sound cans may be distinguished from unsound in a lot of suspicious goods, when the swelling of the ends is not apparent, by the following method:

Boil the cans for an hour, causing the ends of all to swell, then cool, and set aside for eight hours, during which the sound cans will snap back, while the unsound will continue convex, by reason of the fact that the swelling in this case is due to the generation of gas by the bacteria present.

Examination of Gases from Spoiled Cans.—When the tops of blown cans are punctured in the process of opening, an outflow of gas is usually to be noted. Doremus * has studied the character of these gases and



Fig. 119-Apparatus for Collecting Gases from Spoiled Cana. (After Doremus.)

found that when the contents have become putrid, carbon dioxide and hydrogen are the chief gases to be found. Often 60 to 80 cc. of gas may be collected from a can. For the collection of the gases, Doremus

^{*} Jour. Am. Chem. Soc., 19, 1897, p. 733.

uses the device shown in Fig. 119. An adjustable clamp has attached to its upper arm a beveled, hollow, steel needle. A perforated rubber stopper covers the needle and serves as a cushion. connects the needle with the receiver of a eudiometer, both tube and receiver being filled with water or mercury. Either the stop-cock form of eudiometer, as here shown, or the kind with attached leveling-tube may be used. The can is adjusted between the arms of the clamp, and by turning the screw the needle is brought into contact with the top of the can and caused to puncture it, the rubber stopper serving to make a gas-tight joint. The gas passes through the tube into the eudiometer, and its constituents are determined in the usual manner, either by introducing the reagents directly into the eudiometer-tube in the proper order, or by transferring the gases to pipettes.* Hydrogen sulphide is tested for by subjecting a filter-paper moistened with lead acetate solution to the gas. If it turns black, the presence of hydrogen sulphide is indicated.

METALLIC IMPURITIES.

Salts of Lead and Tin are commonly met with in varying amounts in nearly all classes of products put up in tin. The quantity dissolved depends largely on the character of the tin plate used in the manufacture of the can, as well as on how the solder is applied. Much depends also on the nature of the food product and its acidity. much danger was apprehended from the use of the so-called terne plate as a material for cans. This consists of an alloy of lead and tin, coated on iron plate and intended for use as roofing. Sometimes two parts of lead to one part of tin are found in terne plate. Only the better grades of bright tin plate should be used in canning. There is reason to believe that no terne plate is at present used in cans. In 1892 the plating alloy of 47 samples of tin cans in which peas had been put up were examined in the Bureau of Chemistry of the U.S. Department of Agriculture, † and the amount of lead found varied from o to 13 per cent. Only 4 samples were found to exceed 5 per cent, and 24 contained less than I per cent.

The construction of the can should be such that practically no soldered surface is exposed to the contents, the joints being lapped and soldered on the outside. In spite of this, however, it is not unusual to find cans

^{*} See Thorpe's Dictionary of App'd Chem., Vol. 1, pp. 159-161.

[†] Bul. 13, p. 1036.

in which the contents have access to the solder, and it is common to find lumps of solder in the can from the final sealing of the tapped hole in the top. The amount of lead in 24 samples of solder taken from the interior of some of the cans mentioned in the preceding paragraph, was found to vary between the limits of 51 and 65 per cent.*

Action of Fruits and Vegetables on Tin Plate.—The amount of tin dissolved by various canned fruits and vegetables is roughly indicated by the corrosion of the inner surface of the can. A large variety of these canned products have been examined in the laboratory of the Massachu-

Fig. 120.—Interior of Blueberry Cana, Cut Open to Show the Corrosion by Acid of the Fruit Juice.

setts State Board of Health, with a view to determining the quantity of tin contained in solution. The results have shown that though notable traces of tin were found in acid fruits and rhubarb, and large traces in some green vegetables, canned blueberries were found to contain, as a rule, much more tin in solution than any other canned goods examined. It is assumed that the tin was, at least in considerable part, still held in solution by the fruit acids, inasmuch as the metal was found in the filtered juice. In every instance the inner tin lining was found to be extensively corroded, and in some cases it had been almost entirely dissolved off, leaving the underlying iron bare. Fig. 120 shows the appear-

ance of one of these cans, split open to show its inner surfaces. The corrosion is apparent. Eleven samples of canned blueberries, representing seven brands, were examined in 1894 by Worcester, showing an amount of tin in solution (calculated as SnO₂) varying from 0.066 to 0.27 gram per can of 615 cc. capacity.

In 1899 samples of various canned products were examined for lead and tin in the author's laboratory, the results of which are thus summarized:*

	Tin, Grams.	Lead, Grams.	Capacity of Can, cc.
Strawberries,			615
Highest.	.0393	.0004	
Lowest.		.0000	
Raspberries	,		615
Highest	_	.0002	0.3
Lowest		1000.	
Blueberries.	11-3		615
Highest.	1	.0021	013
Lowest	1		
Tomatoes	1	.0004	oro
	L .	••••	950
Highest.		.0004	
Lowest		1000.	é
String beans.			650
Highest.		.0003	
Lowest,	1	.0008	
Peas	1		615
Highest.		.0000	
Lowest		.0001	
Corn		• • • • •	615
Highest	.0101	.0011	
Lowest	. 0045	1000.	
Lima beans		.0004	650
Succotash	0039	.0001	650
Squash,			950
Highest.	- 1793	.0087	
Lowest	-1577	.0003	
Pumpkin		.0010	950
Rhubarb.		.0002	615
Asparagus		.0001	930
Mutton broth		.0001	950
Tomato soup		.0002	370
Salmon		10001	470
Lobster.	1 3-7	.0001	430

A wide range of variation exists in the amount of tin dissolved by various fruits. In the case of pumpkin and squash, for example, the tin dissolved is surprisingly large in quantity, considering the supposed inert nature of these vegetables.

Samples of canned sardines put up in mustard, vinegar, and oil have

^{*} An. Rep. Mass. State Board of Health, 1899, p. 623.

also been examined by the Massachusetts Board, and found to be high in tin. The highest figures showed 0.376 gram (expressed as metallic tin) in a half-pound can. In these cases the corrosion of the interior of the cans was very marked.

Effect of Time on Amount of Tin Dissolved.—A series of experiments was conducted by the author in 1899 * on the action of various fruit acids on tin, with a view to ascertaining, among other facts, whether or not the element of time exerts an appreciable difference in the results.

Samples of various canned fruits and vegetables were titrated for their acidity. It was found that certain samples of canned blueberries, for instance, had an acidity of about one-twentieth normal. In the case of strawberries, the acidity was about one-sixth normal. Canned raspberries were found to be about one-tenth normal in acidity, while the acidity of canned tomatoes varied from one-tenth to one-fourteenth normal. Solutions of one-fifth, one-tenth, and one-fifteenth-normal malic acid, one-tenth and one-fifteenth-normal tartaric acid, one-tenth and onefifteenth-normal citric acid, and one-tenth-normal acetic acid were prepared and sealed in pint glass jars, having about the same capacity as the ordinary-sized tin fruit cans, each jar containing an amount of tin plate equivalent to the interior exposed surface of a can. Solutions thus sealed were kept for three months, six months, and a year, and examined at the end of these respective periods for tin. The results showing the amount of tin found at the end of three months in each case are given in the following list:

ACTION OF FRUIT ACIDS ON TIN IN THREE MONTHS.

Acid.	Grams of Tin in One Pint of Solution.	Acid.	Grams of Tin in One Pint of Solution.
N/5 malic	0.0201	N/15 tartaric	0.0374 0.0236

It was found, in general, that the amount of tin dissolved in three months, as indicated above, was the maximum amount dissolved, or, in other words, with a very few exceptions no additional tin was dissolved by added exposure to the acid for six months, or even a year. The

^{*} An. Rep. Mass. State Board i Health, 1899, p. 624.

amount of tin dissolved was found to vary proportionally with the strength of acid, as would naturally be expected.

Experiments with tenth-normal acetic acid (which was found to be the approximate acidity of the canned sardines mentioned on page 894), sealed in jars with tin plate, as in the case of the fruit acids, and kept for three and six months respectively, showed that in three months 0.0019 gram, and in six months 0.0083 gram of metallic tin had been dissolved, indicating much less vigorous action than that of the same strength of fruit acids, and dissolving less tin than the samples of sardines examined.

Salts of Lead.—While it is a fact that the material of the tin plating usually found in cans is comparatively low in lead, the same is not always true of the metal caps used to cover some of the bottled goods. The French "haricots verts" are usually sold in wide-mouthed bottles, closed by a disk of very soft metal. In one instance this metal cap, which came in contact with the liquid contents of the bottle, was found to contain 93½% of lead. Of the various kinds of bottles in which are sold cheap carbonated drinks known as "pop," one style has a stopper consisting of a metallic button surrounded by a rubber ring. These metallic buttons consist of tin and lead in varying proportions. Inasmuch as the inclosed liquor was usually found to be quite acid in reaction, the danger of prolonged contact with the metallic portion of the stopper is evident.

The following table gives the percentage of lead found in the stoppers of this character, together with the amount of lead contained in the liquor:*

Character of Sample.	Per Cent of Lead in Stopper.	Amount of Lead in Contents of Bottle in Milli- grams.†
Blood orange	50.7	0.31
Birch beer	35.0	Large trace
Ginger	32.2	0.40
Strawberry A	8.8	0.20
Strawberry B	6.5	0.30
Sarsaparilla A	8.5	0.19
Sarsaparilla B	3.5	0.17
Lemon	7.5	0.27
Maximum	50.3	1.05
Minimum	3.8	10.0

† Capacity of bottle about } pint.

Besides the above tabulated samples, twenty were found with stoppers containing less than 3% of lead. While the amount of lead found in the

^{*} An. Rep. Mass. State Board of Health, 1897, p. 571.

contents of the bottles was in no case very large, it was enough to condemn the use of lead in the manufacture of such stoppers. That the amounts of lead found in the contents of the bottles vary quite irrespective of the percentage of lead in their stoppers, may be ascribed to various causes, such as the difference in the acidity of the liquors, and the length of time that the liquor has been in contact with the stopper. Furthermore, the more soluble metal of an alloy is attacked by an acid with an energy which is not proportional to the percentage of that metal in the alloy.

Salts of Zinc.—The presence of zinc salts in canned foods is largely accidental, and is generally due either to the contact of the acid fruits and vegetables with galvanized iron in the canneries, to the occasional use of brass vessels, or to the zinc chloride used as a soldering fluid. Hilgard and Colby * have examined empty tin cans fresh from the manufacturer, and found zinc chloride in notable quantity in the seams, and especially in the empty space of the lap at the bottom of the can, where it could easily be acted on by the contents. The average amount of soluble zinc chloride found in the "lap" alone amounted to three-fourths of a grain per can. It was furthermore ascertained that it was not the practice of canners to wash the cans before packing, so that zinc present in canned goods may thus readily be accounted for.

Zinc chloride is commonly used in machine soldering, but should be displaced by rosin.

Hilgard and Colby found in some spoiled cans of asparagus, where the acidity was unusually high, an average of 6.3 grains of zinc chloride per large can.

Zinc salts are said to have been used for greening peas, but their use for this purpose is not common. Zinc chloride is the salt used, and a natural yellowish-green tint is imparted when properly applied. The process has been kept secret.

Salts of Copper.—While copper in canned goods is sometimes accidental, its presence being due to the use of copper or brass vessels in the canneries, its chief interest to the food analyst lies in the use of copper sulphate for greening peas and other vegetables. The artificial greening of vegetables is much more commonly practiced in France than in the United States.

French canners of peas, beans, Brussels sprouts, etc., are frequently so lavish in the use of sulphate of copper that the goods as found on our

^{*} Rep. Cal. Agric. Exp. Sta., 1807-8, p. 150.

markets can in some cases hardly be said to resemble the freshly cooked products in color. Oftentimes, indeed, they possess such a deep green as to be positively distasteful to the average American palate, though evidently this unnatural hue is craved in Europe. The use of copper in such foods is often rendered apparent by the most cursory examination.

In this country, when copper is used, smaller quantities are usually employed, with an attempt to imitate more closely the color of the natural product.

Complaint in court for this form of adulteration under the general food law as it exists in most states would naturally be brought under one of two clauses:

1st. As being colored, whereby the product appears of greater value than it really is, or

2d. As containing an ingredient injurious to health.

An ingenious claim is sometimes advanced by the defendant in opposition to clause 1, to the effect that copper sulphate is added, not to give an artificial green color, but to preserve the original green of the chlorophyl or natural color of the fresh peas,* so that it will not be destroyed by subsequent boiling.

This point was argued in a strongly contested court case brought in Massachusetts for copper in French peas.

As Worcester ‡ has shown, the fallacy of this argument can be easily demonstrated. If it were true that the copper acts as a preservative of the chlorophyl, a pure extract of chlorophyl should, by the addition of copper sulphate, be prevented from destruction on boiling, and again, on once destroying the color of the chlorophyl by boiling, it would be impossible to restore it by further boiling it with copper sulphate.

As a matter of fact, if an extract of chlorophyl is boiled with a dilute solution of copper sulphate, its color is at once destroyed, and a brown precipitate is thrown down. On the other hand, if yellow or white peas or beans devoid of chlorophyl are boiled with copper sulphate, they are colored green, the depth of color depending on the strength of the copper solution. When peas or other vegetables are thus colored, very little copper is found, as a rule, in the liquid contents of the can, but the copper is chiefly confined to the solid portions. Green compounds are produced

^{*} The term used by the French to describe this process is reverdissage or "regreening."

[†] An. Rep. Mass. State Board of Health, 1892, p. 605.

t Loc cit., supra, p. 641.

by boiling albumins with copper salts, due to the fermation of albuminate, or in the case of peas, leguminate of copper. Harrington * states that it is possible to color eggs an intense green by boiling with copper sulphate.

Examination of a large number of brands of canned vegetables greened by copper, as bought in Massachusetts, showed that the amount used varied from a trace to 2.75 grams per can, calculated as copper sulphate. In justice to the consumer, who may be cautious about taking into his system copper salts, as well as to those who are indifferent to their use, it is no more than fair that all cans should have a label, plainly stating the quantity present. In the Massachusetts market, labels like the following are not uncommon: "This package of French Vegetables contains an equivalent of Metallic Copper not exceeding three-quarters of a grain."

Copper as a coloring matter has been most commonly found in peas, beans, and Brussels sprouts. Copper salts in minute quantity have been found in Massachusetts in canned tomatoes, clams, and squash, as well as in pickles.

Salts of Nickel.—Sulphate of nickel has been employed instead of sulphate of copper for greening vegetables. According to Harrington † 0.25 gram of nickelous sulphate per kilogram of peas is used. The peas or other vegetables are boiled in a solution of the salt, made slightly alkaline with ammonia.

Toxic Effects of Metallic Salts.—Divergence of opinion is so great as to the toxic effects of salts of the heavy metals on the human system, when present in the small amounts commonly found in food products, that it is extremely difficult to maintain a complaint in court based entirely on the harmful effects of these salts. Since the question is one for the toxicologist or physiological chemist rather than the analyst to settle, it will not be discussed here at length; suffice it to say that a large number of experiments on human beings will undoubtedly have to be tried, before the necessary data will be at hand on which to base a really intelligent opinion. The same general difficulties are met with here as one encounters in the matter of determining the definite effects of antiseptics in food, of alum in baking-powder, etc.

Determination of Lead in Tin Alloy.—Method of Paris Municipal Laboratory.‡—The material, if soft, is hammered into a thin plate, and

^{*} Practical Hygiene, p. 203.

[†] *Ibid.*, p. 205.

[‡] Analyse des Matières Alimentaires et Recherche de leurs Falsifications, 1894, p. 605.

2½ grams are weighed out, transferred to a 250-cc. flask, and dissolved in 7 to 8 cc. of concentrated nitric acid. Evaporate to dryness on the sand-bath, add 10 drops of nitric acid and 50 cc. of boiling water, cool, and make up to 250 cc. with water. Let the residue settle and pour off through a filter 100 cc. of the clear, supernatant liquid, corresponding to 1 gram of the material. This contains the lead, while the tin is left behind in the residue, together with antimony if present.

Add 10 cc. of a standard solution of potassium bichromate (7.13 grams to the liter) and shake. Each cubic centimeter of this standard solution is sufficient to precipitate 0.01 gram of lead. Allow the lead chromate formed to settle, and, if the solution is colorless, add 10 cc. more of the bichromate, or sufficient to be present in excess, as indicated by the yellow color. Filter, wash, and titrate the excess of bichromate with a standard iron solution, containing 57 grams of the double sulphate of iron and ammonia and 125 grams of sulphuric acid per liter. This iron solution should be kept under a layer of petroleum, and standardized against the potassium bichromate before use.

Add, drop by drop, the iron solution to that containing the excess of bichromate. The color of the latter passes from pale green to bright green, when the chromate is completely reduced. Determine the endpoint with a freshly prepared dilute solution of potassium ferricyanide, a drop of which is placed on a porcelain plate or tile in contact with a little of the solution titrated. A blue color is produced when the iron is present in excess. If the standard iron and bichromate solutions exactly correspond, I cc. of the iron solution is equivalent to 1% of lead, but the latter solution is usually a little weak.

If n= number of cubic centimeters of iron solution necessary to reduce 10 cc. of the standard bichromate,

I cc. of the iron solution =
$$\frac{10}{n}$$
.

If, now, r=number of cubic centimeters of iron solution necessary to reduce the excess of bichromate in the determination, and s=number of cubic centimeters of bichromate used,

$$s - \frac{10}{n}r = per cent of lead in the alloy.$$

Separation and Determination of Tin, Copper, Lead, and Zinc in Canned Goods. — Munson's Method.*— The contents of the can are

^{*} U. S. Dept. of Agric., Bur. of Chem., Bul. 65, p. 52.

first evaporated to dryness, and from 10 to 15 cc. of concentrated sulphuric acid or enough to carbonize are added to the dry residue contained in a porcelain evaporating-dish, which is very gently heated over the flame till foaming ceases. Then ignite to an ash in a muffle, or carefully over the free flame, using a little nitric acid, if necessary, for oxidation of the organic matter. Add 20 cc. of dilute hydrochloric acid, and evaporate over the water-bath to dryness. Wash the residue into a beaker, slightly acidify with hydrochloric acid, and saturate with hydrogen sulphide without previous filtration. Heat the beaker on the water-bath, and pass the contents through a filter. Wash the precipitate, which contains sulphides of tin, lead, and copper, if these metals are present, while if there is zinc, it is contained in the filtrate. The precipitate is fused with sodium hydroxide in a silver crucible for half an hour, to increase the solubility of the tin, which would otherwise be dissolved with difficulty. The fusion is boiled up with hot water, acidulated with hydrochloric acid, and transferred without filtering to a beaker, in which hydrogen sulphide is added to saturation. This precipitates the sulphides of tin, lead, and copper (if these metals are present). The sulphide precipitate is collected on a filter, and thoroughly washed with hot water, the washings being rejected. Pass through the filter several portions of boiling ammonium sulphide, using about 50 cc. in all, or till all the tin is dissolved. Precipitate the tin from the combined filtrate with hydrochloric acid, filter, wash, ignite, and weigh as stannic oxide.

The residue left on the filter, after dissolving out the tin sulphide, is then dissolved by treatment with nitric acid, which is filtered, and to the filtrate and washings ammonia is added nearly to the point of neutralization. Then add ammonium acetate. Filter off any precipitate of iron that may be formed. The filtrate is divided into two portions for determination of copper and lead. If lead is absent, determine the copper by titration with potassium cyanide * or electrolytically (p. 608). Copper is rarely present in sufficient amount to be determined, unless used for greening the vegetables. If notable quantities of lead are present, the solution is made acid with acetic, and the lead precipitated therefrom with potassium chromate, collected on a tared filter, washed with water acidified with acetic acid, dried at 100° C., and weighed as lead chromate. Or determine the lead by color-tests, as on page 902.

For the determination of zinc, the filtrate from the first hydrogensulphide residue is evaporated to a volume of about 60 cc., and treated

^{*} Sutton, Volumetric Analysis, 8th ed., p. 204.

with bromine water to oxidize the iron, as well as any excess of hydrogen sulphide remaining, the excess of bromine is then boiled off, and a few drops of concentrated ferric chloride added, to make the solution distinctly yellow, if not already so. Nearly neutralize with ammonia, and precipitate the iron with ammonium acetate. Filter, wash, acidify the filtrate with acetic acid, and precipitate the zinc with hydrogen sulphide. Filter, wash, ignite, and weigh as zinc oxide.

The metals may be determined separately, as follows:

Determination of Tin.*—Evaporate the contents of the can to dryness, and ignite in porcelain. Fuse the ash with sodium hydroxide in a silver crucible, boil the fusion with several portions of water acidulated with hydrochloric acid, filter, and precipitate the tin from the acid solution with hydrogen sulphide. Dissolve the washed precipitate in ammonium sulphide, filter, and deposit the tin directly from this solution by electrolysis in the platinum dish which contains it, using a current of 0.5 ampere and the electrolytic apparatus described on page 608.

Determination of Lead, especially applicable if lead is present in small amounts only. Boil the sulphated ash of the contents of the can (obtained as on page 901) with a solution of ammonium acetate, having an excess of ammonia. The tin, zinc, and iron remain insoluble, while the copper and lead are dissolved. Filter, wash, and add a few drops of potassium cyanide to the filtrate, to prevent precipitation of copper when hydrogen sulphide is subsequently added. If the solution exceeds 40 cc., concentrate to that amount by evaporation, and transfer to a 50-cc. Nessler tube. Add hydrogen sulphide water, and make up to the mark. Compare the brown color imparted by the lead sulphide, with the colors obtained by treating with hydrogen sulphide water in Nessler tubes various measured amounts of a standard solution of lead acetate, made alkaline with ammonia.

Determination of Copper.—(1) Electrolytically.—Ash the contents of the can as on page 901. Wet the ash with concentrated nitric acid, add water, and boil Then make strongly alkaline with ammonia and filter. Unless the filtrate is colored blue, copper is absent. Transfer the filtrate to a bright tared platinum dish of 100-cc. capacity, neutralize with concentrated nitric acid, and add about 2 cc. in excess. Nearly fill the dish with water, and electrolyze with the apparatus described on page 608, using a current of about 0.3 of an ampere.

^{*} Hilger u. Laband, Zeits. für Untersuch. Nahr. u. Genuss., II, p. 795; An. Rep. Mass. State Board of Health, 1899, p. 625.

(2) Colorime!rically.—This method is especially applicable for small amounts of copper. The blue-colored ammoniacal solution of the ash, filtered as in (1), is transferred to a Nessler tube, and its color matched against the colors of a series of measured amounts of an ammoniacal standard solution of copper sulphate

Determination of Nickel.—Boil the ash with water slightly acidified with hydrochloric acid, and without filtering, saturate with hydrogen sulphide, thus precipitating out any copper, tin, or lead. Filter and wash. Zinc and nickel, if present, are in the filtrate. Boil the filtrate to expel the hydrogen sulphide, and add sodium carbonate till slightly alkaline. Add acetic acid without filtering till the precipitate produced by the alkaline carbonate is dissolved, and then add a considerable excess of acetic acid. The zinc is precipitated by passing hydrogen sulphide through the cold dilute solution, while the nickel is held in solution by the large excess of acetic acid. Filter, and wash with hydrogen sulphide water, to which a little ammonium acetate has been added.

Make the filtrate alkaline with ammonia, precipitate the nickel with ammonium sulphide, filter, wash, ignite, and weigh as nickelous oxide.

ANTISEPTICS IN CANNED FOODS.

No class of food products stands so little in need of these added substances to arrest fermentation as canned foods, if properly prepared; but, as a matter of fact, the use of antiseptics in this connection is still practiced. Prolonged heating for a sufficient length of time to perfectly sterilize the contents of a can is in some cases more or less detrimental to the appearance of the product, so that for this reason, as well as to save time in "processing," and furthermore to increase the keeping qualities of the goods after opening, many manufacturers resort to the use of artificial chemical preservatives. So long as any well-founded prejudice against preservatives exists, their use in canned or bottled foods should be unequivocally condemned, unless the cans or packages are distinctly labeled with the nature of the preservative and the extent to which it is employed.

As in other foods, discrimination as to locality is apparently used on the part of manufacturers in shipping canned goods containing added preservatives, so that, as a matter of fact, in states where it is well understood that a vigilant enforcement of the pure-food law prevails, we do not find as high a percentage of canned foods with preservatives as in other states. With the national law, preservatives are now less prevalent. Preservatives commonly employed in canned goods are salicylic, benzoic, and sulphurous acids, though the other familiar antiseptic agents may be used. In such foods as canned corn, while the purpose of sulphurous acid may be in part as a preservative, the primary object for its use is undoubtedly to bleach or whiten the product.

The Bleaching of Corn by artificial means before canning is usually accomplished by boiling the corn with sulphite of soda, thus giving to the product an unnaturally white color. The practice seems to have been more in vogue ten years ago than at present, the popular taste now apparently preferring the natural rich yellow of fresh corn.

Saccharin is claimed to possess antiseptic powers and is used in canned goods, but its primary purpose is as a sweetener.

Beta-naphthol is also said to be used as a preservative in canned goods, but has not been found by the author in any samples that have come to him for analysis.

Detection of Preservatives.—Tests for salicylic or benzoic acid are most readily made in the residue from an ether extract of a portion of the acidified contents of the can or package, while formaldehyde and sulphurous acid are tested for in the earlier portions of the distillate, obtained by distilling a mixture of the acidified contents in water.

If it is desired to systematically test for the various preservatives in the same sample, a convenient method of procedure is as follows:

Thoroughly mix 50 grams of the pulped sample with water in a 250-cc. graduated flask, make distinctly acid with dilute phosphoric acid, and fill to the mark with water. Transfer to a distilling-flask, and subject to distillation in a glycerin- or paraffin-bath, whereby the temperature is raised near the end of the distillation considerably above 100° C.

Remove the first 30 cc. of the distillate, and divide into three equal portions, which are to be tested for formaldehyde, sulphurous acid, and beta-naphthol by the usual tests for these preservatives.

Continue the distillation till the residue in the flask is nearly dry, and transfer the remaining or larger portion of the distillate to a large separatory funnel. Acidify with dilute sulphuric or hydrochloric acid, and extract with ether or chloroform.

Divide the ether or chloroform extract into three portions in as many evaporating-dishes, evaporate to dryness at low temperature, and make the appropriate tests on the three residues for salicylic acid, benzoic acid, and saccharin, as given in Chapters XVIII and XIX.

The residue left in the flask is then washed out and incinerated, and the ash examined for boric acid.

"SOAKED GOODS."

It has become quite common, especially in the case of peas, beans, and corn, to utilize for canning purposes those that have grown old and dried, after soaking them for a long time. The presence of soaked peas in the market is generally more common in years when there is a scarcity in the pea crop. By the process of soaking, dried and matured field corn may be softened to such an extent as to be substituted for green or sweet corn in the canned product. These goods, frequently sold at a very low price, under some such tempting name as "Choice Early June Peas," are entirely devoid of that succulent property so highly prized in the fresh goods, and are altogether so inferior in quality that their sale may justly be considered as fraudulent, unless their character is specified. In some states the law provides that such a product, to be legally sold, shall have plainly marked on the label of the can the words "Soaked Goods" in letters of prescribed size.

Detection.—Methods of detecting soaked goods are distinctly physical rather than chemical. The appearance and taste of the goods furnish in most cases an unmistakable clue to their nature. Soaked goods are entirely lacking in juiciness, and in the flavors so characteristic of the various vegetables, when gathered and canned before becoming dry. The process of soaking is also said to develop the growth of the rudimentary stem of the embryo in the dried pea and bean. Peas and beans of the soaked variety are almost entirely lacking in the green color of the fresh vegetables, unless the color has been artificially supplied.

In all cases it will be found that the solid grains or kernels of the peas, beans, and corn that have once been dried, though softened by the process of soaking, have much less water than the grains of the corresponding vegetables that were gathered while still soft and succulent.

KETCHUPS AND TABLE SAUCES.

These preparations vary widely in their character and composition, and, in the absence of standards fixed by law for each particular mixture, almost any food product may be included in its make-up, without laying it open to the charge of being adulterated. At the same time, in this class of condiments it is naturally expected that the ingredients used all have some food value, and in addition possess a certain degree of pungency

	Number of Analyses	Refuse.	Water.	Protein.	Pat.	Total Carbo- hydrates	Ash.	Fuel Value per Pound.
Tomato ketchup	2		82.8	1.5	.2	12.3	3.2	265
Horseradish	2		86.4	1.4	.2	10.5	ĭ.5	230
Olives, green:			_			1		١.
Edible portion	I		58.0	1.1	27.6	11.6	1.7	1,400
As purchased	I	27.0	42.3	.8	20.2	8.5	1.2	1,025
Olives, ripe:	1							
Edible portion	1		64.7	1.7	25.9	4-3	3-4	1,205
As purchased	1	19.0	52.4	1.4	21.0	3.5	2.7	975
Cucumber pickles	3		92.9	-5	-3	2.7	3.6	70
Mixed pickles	I		93.8	1.1	-4	4.0	-7	110
Spiced pickles	I		77-1	-4	- I	20.7	1.7	395

CHEMICAL COMPOSITION OF KETCHUP, PICKLES, AND RELISHES.*

* U. S. Dept. of Agric., Office of Exp. Sta., Bul. 28, p. 70.

or distinctive flavor. In other words, inert materials used simply as "fillers" are, to say the least, out of place, even though they are not actually adulterants.

Tomato Ketchup of the U. S. Standards, consists of the strained pulp of boiled, fresh, ripe tomato mixed with various spices, either with or without the addition of sugar and vinegar.

The ketchup of the housewife is made from varying recipes, all based on the above method of procedure; and while the commercial bottled ketchups should be made from materials quite as pure, it is often true, especially in the cheaper varieties, that the skins and refuse of tomatocanning factories, pumpkin pulp, or apple pulp, form the basis of the product. Even with the use of these materials, when properly prepared, and before advanced fermentation has set in, with clean methods of handling, the product may not be unwholesome. It is, however, sometimes the practice to allow the refuse and skins to accumulate through a whole tomato-canning season, storing them all in large vats, and working them up, after they have become badly fermented, for "fresh tomato ketchup." It is largely for this reason that antiseptics and coloring matters are so commonly employed in ketchup.† Salicylic acid, formerly much used as a preservative of ketchup, has in the past few years given place to benzoate of soda.

Bitting has shown that by using sound tomatoes and exercising

[†] The writer has in his possession a circular from an Indiana commission merchant, advertising for sale tomato pulp of some twelve different grades for ketchup. Among them are listed the following: "100 bbls. of old goods, made partly from whole stock and partly waste, boiled down nearly to ketchup thickness; has preservaline in it; fine goods, but some of it is fermented; packed in good oak whiskey and wine barrels. Price \$2.00 per bbl."

proper care in the process of manufacture, ketchup can be kept without a preservative.* Manufacturers are themselves corroborating this.

Coloring of Tomato Ketchup.—The practice of adding artificial dyestuffs to ketchup is decreasing. Indeed the very brilliant scarlet and crimson hues sometimes given to the bottled ketchups on the market in no wise resemble the natural dull-red or brown color of the pure homemade article, in which the bright color of the fruit pulp is modified by the mixture of spices with which it is cooked. It is doubtless true that many manufacturers employ such inferior materials that unless some dyestuffs were added the result would be most unappetizing in appearance. Out of ninety-five samples of tomato ketchup examined in 1901 in Connecticut all but fifteen contained coal-tar colors.†

Walnut Ketchup.—This is made up in a somewhat similar manner to tomato ketchup, excepting that instead of tomatoes, soft young walnuts are crushed and used as a basis.

Chili Sauce is made up of a pulped mixture of tomatoes, red peppers, onions, vinegar, and various spices, differing from ketchup in that it contains the seeds and is not strained. In consistency it is heavier than ketchup. It is colored and preserved in much the same manner as ketchup.

Table Sauces are composed of a large variety of materials of a more or less pronounced flavor or pungency, combined in a liquid preparation usually of a more thin or watery consistency than the ketchups. The materials employed include mushrooms, onions, garlic, ground anchovies, tamarinds, spices, coriander and cardamom seeds, walnuts, vinegar, molasses, and even assafætida. These bottled preparations are very rarely colored except with caramel, but sometimes contain antiseptics, especially salicylic and benzoic acids.

The Acidity of ketchups and table sauces furnishes a ready means of comparison between different varieties, and is conveniently expressed in terms of acetic acid.

To determine the acidity, titrate I gram of the diluted sample with tenth-normal sodium hydroxide, using phenolphthalein as an indicator.

[&]quot;225 bbls. new goods, made from waste; has bensoate of soda in it, packed in uncharred whiskey and wine barrels at \$3.00 per bbl. net cash." "300 bbls. old goods, partly whole stock, partly waste, has salicylic acid in it; nice goods, etc. Price \$2.00 per bbl." "400 bbls. new goods, Jersey style; solid and good red color, fine quality. Price \$3.00 per bbl." With prices as low as the above quotations, it is difficult to see how a cheaper basis for ketchup stock than the above could be supplied. Even the pulp of pumpkin and of other inert vegetables, alleged to be used as adulterants, would hardly be furnished so cheaply.

^{*}U. S. Dept. of Agric., Bur. of Chem., Bul. 119.

[†] An. Rep. Conn. Exp. Sta., 1901.

Each cubic centimeter of the alkali corresponds to 0.006 gram of acetic acid.

Winton and Ogden * have found the acidity of tomato ketchups examined by them to vary between the limits of 0.60 and 2.20 per cent, calculated as acetic acid, Chili sauce from 0.80 to 1.80 per cent, and various table sauces from 1.40 to 1.60 per cent.

Examination of Table Sauces and Ketchups for Preservatives.— Extract a portion of the acid sample with ether or chloroform, which removes salicylic or benzoic acid or saccharin. If the sample is of thin or watery consistency, like most table sauces, the extraction can in most cases be readily effected in a separatory funnel, chloroform being in this case most convenient, since it sinks to the bottom. If ketchup or other thick syrupy substance is to be examined, it is almost impossible when shaking with ether or chloroform to avoid the formation of an annoying emulsion, which it is difficult to break up. For this reason the author prefers, in the case of ketchups and similar viscous fluids, to separate the extract by means of a centrifuge of the style shown in Fig. 11. A portion of the acid sample, say 75 cc., is shaken violently in a corked flask with 25 to 40 cc. of ether, and the mixture, usually in the form of an emulsion, is poured into two of the centrifuge tubes, so that they contain equal amounts and balance each other. They are then corked, placed in the shields of the centrifuge, and whirled from two to four minutes, or until the emulsion is broken up. At the end of this time it is usually found that the mixture is separated into three layers: first a watery layer at the bottom of the tube, then an almost solid layer of the viscous material above it, and finally the clear ether extract at the As a rule the separation is so complete that the tube may be inverted, and every drop of the clear ether layer may be decanted without filtration, so firmly does the solid middle layer hold in place in the tube. Indeed, a vigorous shake is usually necessary to dislodge it.

If saccharin, as well as salicylic and benzoic acids are all to be looked for, the ether extract is divided into three portions, in as many evaporating-dishes, and the dried residue tested in the regular manner for the above substances.

Examination of Ketchups for Colors.—Proceed as in the case of jellies and jams.

^{*}An. Rep. Conn. Exp. Sta., 1901, p. 137.

PICKLES.

A large variety of vegetables and fruits are preserved in the form of pickles in vinegar, either with or without spices, and kept in wooden pails, stoneware pots, kegs, or sealed, wide-mouthed bottles. The containers are not of necessity air-tight. The commoner vegetables are usually pickled without cooking, while with fruits, as in the case of peaches, pears, gooseberries, etc., they are usually cooked, or at least heated.

Cucumber Pickles are the most common, and are prepared by soaking the fresh cucumbers in strong salt brine. They are then dried on frames, and afterwards treated with boiling vinegar, to which spices may or may not be added. Other vegetables pickled in similar manner, either separately or in mixture with cucumbers to form "mixed pickles" or "gherkins," are cauliflower, bean pods, white cabbage, young walnuts, and onions.

Such soft vegetables as young podded beans and beets are not treated with brine, but, after soaking in water, are directly treated with vinegar. The vinegar used for the finest pickling is of the cider, wine, or malt variety. Cheaper varieties of pickles are put up in "white wine" or spirit vinegar.

Mustard Pickles.—These differ from plain vinegar pickles in the character of the preserving medium, which in this case consists of a mixture of mustard and spices with the vinegar to form a thin paste.

Piccalilli consists of a mixture in vinegar of various chopped vegetables, such as cucumbers, cauliflower, green pickles, onions, green tomatoes, and various spices.

Olives for pickling are picked before they have fully ripened, and the inherent bitter taste is removed by soaking in a solution of potash and lime. This is replaced by cold water, and finally the olives are transferred to the medium in which they are bottled, which consists of salt brine, either with or without flavoring. The flavoring materials employed consist of such substances as fennel, coriander, laurel leaves, and occasionally vinegar.

Capers.—These are the flower buds of the shrub Capparis spinosa, which are pickled in vinegar. Nasturtium seeds, when similarly pickled, possess a flavor much resembling capers, but their substitution for capers could readily be detected by their distinctive appearance, even if colored.

Adulteration of Pickles.—Green pickles, such as cucumbers, are

not uncommonly colored artificially by copper salts, either through the addition of copper sulphate, as in the greening of peas, or by the use of copper vessels. This artificial greening is to be looked for also in such products as capers and olives.

For methods of detection and estimation of copper, see page 902. Pickles may be greened by boiling with much less harmful substances than copper salts, such, for example, as grape leaves, spinach, or parsley.

Free Sulphuric Acid has been found in a number of cases in the vinegar of pickles bought on the Massachusetts market. A pronounced test for chloride with nitrate of silver should not be attributed to free hydrochloric acid, since it may be and probably is due to the salt from the brine in which the pickles have been treated.

Alum is sometimes added to the salt solution to produce hardness and crispness in pickles. A number of samples of cucumber pickles have been found by the author to contain alum. For its detection, fuse the ash of the pickles, if free from copper, in a platinum dish with sodium carbonate, extract with boiling water, filter, and add ammonium chloride. A flocculent precipitate shows alum.

Sodium Benzoate and Saccharine are frequently used in sweet pickles. Horseradish.—This condiment is prepared by grating the root of the perennial herb Nasturtium armoricia, and preserving in vinegar. It is very pungent and aromatic when first prepared, but by exposure to light and air quickly loses strength. Turnip, an occasional adulterant of grated horseradish, is best detected by the microscope.

JAMS AND JELLIES.

Jams or marmalades are prepared from the pulp of fruits, and jellies from the fruit juices. Both jams and jellies, to be considered of the highest degree of purity, should contain nothing but the fruit pulp or juice named on the label, mixed with pure cane sugar, and, in the case of jams, the further addition of spices and flavoring materials is permissible.

For the manufacture of jam, the washed fruit, if of the kernel variety, is peeled, freed from cores, and sliced; if berries, they are simply stemmed; if stone fruits, they are peeled, freed from stones, and quartered. The material, properly prepared, is cooked with as much water as is necessary for boiling, and with the addition of an amount of sugar varying with different manufacturers. Some prefer to use equal parts of sugar and fruit, others one part sugar to two parts fruit.

In the case of jelly, the fruit is cooked in a small amount of water

till soft, transferred to a bag or press, and the juice allowed to flow out spontaneously, or is squeezed out under pressure, according to the grade of jelly desired, the clearest and finest varieties being made from the juice that flows out naturally. This juice is then evaporated down with the addition of sugar to a density of from 30° to 32° Bé., which is of the proper consistency to form a perfect jelly product after cooling, and, while still hot, is poured into the tumblers in which it is to be kept. Here, as in the case of jams, the amount of sugar varies, some using pound for pound, and others only half as much sugar as fruit. Some manufacturers clarify their jellies by mixing with the juice, while boiling, elutriated chalk, using a teaspoonful to each quart of juice. The impurities come to the surface with the chalk as a scum, and are skimmed off. This clarifying process is somewhat analogous to the defecation of sugar juices with lime, and is commonly carried out with apple jelly.

The "jellying" or gelatinizing of the final product is due to the presence in the fruit juice of pectin, or so-called vegetable jelly (C₃₂H₄₀O₂₈4H₂O); see page 276.

The high content of added sugar in jelly, once thought to be essential for keeping it, is now no longer considered necessary, and much less sugar is at present added than formerly. The finest grade of apple jelly, for instance, is made without any added sugar whatever.

In making the better grades of apple jelly, apple juice fresh from the press is run directly into the boiler or evaporator before any fermentation has ensued, and gelatinized by concentration. If boiled cider is wanted instead of jelly, it is drawn off at an earlier stage than in the case of apple jelly.

Composition of Known-purity Jellies and Jams.—In the tables on pp. 912 and 913, due to Tolman, Munson, and Bigelow,* are given results reached on the examination of the pure finished products, as well as on pure fruit juices and pulp used in their manufacture.

Adulteration of Jams and Jellies.—As a matter of fact, a small percentage of these products sold in the United States are honest prototypes of the home-made jams and jellies, which consist exclusively of the fruit specified on the label, in mixture with pure cane sugar. If we accept as a standard the product of the housewife, fully 90% of the commercial brands of these preparations would be found wanting. So great is the demand for cheap sweets of this variety, that the market is flooded with them at eight and ten cents per half-pound jar, when in reality abso-

^{*} Jour. Am. Chem. Soc. (1901), pp. 349-351.

TABLE I.-JUICE.

			Total			Sugara.	4			Polarization.	ا ا
Description of Sample.	Total Solids, Per Cent.	Ash, Per Cent,	Acids Calcu- lated as H.SO.	Proteids (NX 6 as), Per Cent.	Reducing Sugar, Per Cent.	Cane Sugar Added, Per Cent.	Cane Sugar Found, Per Cent.	Added Cane Sugar Inverted. Per Cent.	Direct at	Invert at	Invertat 86° C.
Apple (fall pippin)	7.95	0.47	0.627	0.543	8:4		1.18		-3.0	-4.6	-2.9
Blackberry	.54	0.52	0.978	0.320	4-34	•	0.0		-1.5	-1.6	0.1
Crab apple		0.30	0.372	0.075	2.56		1.03		0.I-	-2.4	1:1-
Grape (fox)		0.49	1.686		2.79	:	0.37	:	8.01	-1.3	1.1
Grape (Ives seedling)		0.57	0.902	0.237	5.10		8.0		-I.2	12.4	9.0
Huckleberry	16.33	0.40	0.454		11.21	1 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4	68.0		13.3	4.4	6.0-
Orange (Florida navel)	۰. و	0.30	0.297	0.581	1.52	:	2.29	:	+1.8	-1.3	0.0
Peach	8.8	0-45		0.218		:	4.59		+4.0	-2.2	1.0-
Pear (Bartlett).	11.65	0.45	0.345	0.087	5.87		1.18		4.4	-6.4	0.4-
Pineapple		0.45	0.588	0.368	2.74		8.96	:::::::::::::::::::::::::::::::::::::::	+8.4	13.7	-1.1
4	_	0.77	:	0.350			4-73	:	+4.1	12.3	10-7
	12-72	0 63		0.431	4.86	:	0.51		+3.0	+1.3	+3.4
Plum (wild fox)	_	0.64	1.576	0.137	2-87		2.81	:	+I-4	12.4	00 m
Mixed fruit	_	0.32	0.612	0.150	2.68		0.59		0.1	-1.8	6.0-
				TABLE II TELLY	TELLY						

9.0 0.0 0.0 +0.2 4.0-10.0 9.0 9.1-0 0 0.0 9.0-119.0 118.9 -20.6 -20°I 24.0 36.17 18.20 58.88 49.33 47.54 11.16 11.16 7.33 46.97 66.18 40-38 33.04 23.04 33.05 30.52 30.52 55.50 55.50 22.67 22.67 25.67 25.67 25.67 25.67 51.76 54.89 56.39 65.39 65.39 65.39 65.39 65.39 65.39 65.39 65.39 65.39 65.39 65.39 65.39 65.39 65.39 65.39 24.25 24.27 24.27 24.27 25.29 25.20 0.243 0.1375 0.1375 0.175 0.175 0.350 0.350 0.000 0.17 0.328 0.475 0.171 0.524 0.245 0.171 0.245 0.352 59.18 539.58 63.28 63.00 69.58 69.13 76.34 545.56 73.01 Huckleberry.
Orange (Florida navel).
Peach.
Pear (Bartlett). Pineapple Plum (wild fox).
Plum (wild fox) boiled down. Plum (Damson). Mixed fruit...... Crab apple. Grape (Ives seedling). Blackberry Apple (fall pippin).

The "juice" was prepared by cooking the fruit till soft, after the addition of sufficient water to prevent scorching, and straining through a jelly bag,

*The composition here given is not that of the original fruit, but of the pulped mass used in the preparation of jame.

TABLE III.-FRUIT.

			Total		<u> </u>	Sug	Sugars.		"	Polarization,	
Description of Sample.	Total Solids, Per Cent.	Ash, Per Cent,	Acids Calcu- lated as HrSO4, Per Cent.	Proteids (N×6,25), Per Cent	Reducing Sugars, Per Cent.	Cane Sugar Added, Per Cent.	Cane Sugar Found, Per Cent.	Added Cane Sugar Inverted. Per Cent.	Direct at 18° C.	Invertat	Invert at
Apple (fall pippin). Blackberry Crab apple. Grape (Ives seedling). Orange (Florida navel).	9.25 9.62 14.34 12.50 13.11	0.0000 0.	0.499 0.916 0.705 0.686 0.392	0.725 0.418 0.985 0.056	5.67 5.67 5.68 5.64 5.13		1.03		++ ++	1 1 1 1 1 1 1 1 1 1	# 0 0 0 0 0 1 1 1 1
				TABLE IVJAM.	/.—JAM.				:		
	63.22 61.82 61.80 61.80 61.82 61.52 61.52 62.43	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	0.28a 0.715 0.715 0.744 0.163 0.163 1.355	0.175 0.493 0.525 0.944 0.312 0.312	25.52 14.80 33.06 33.44 13.20 13.20 13.20 13.20 28.23	\$1.31 36.09 36.09 56.13 66.13 47.75 47.86	29.22 23.24 24.23 24.24 25.24 25.24 25.25 25 25 25 25 25 25 25 25 25 25 25 25 2	43.22 34.68 35.46 92.96 73.38 18.87 22.90 74.42	4+++++++ 6427 428 842 50420 428 842 50420 5042 50420 br>50420 50420 50420 50420 50420 50420 50420 50420 50420 50420	0.0000000000000000000000000000000000000	++ + ++++ 4 = 0 4 0 4 = 0 = 0 8 0 0 4 0 0 0 4 4 0

lutely pure goods cannot be produced at much less than twice that amount.

The cheap substitutes are made up largely of apple juice and commercial glucose, sometimes containing no fruit whatever of the kind specified on the label. Sometimes an attempt is made to imitate the flavor by the addition of artificial fruit essences, but more often the same apple-glucose stock mixture of jelly, put out under a particular brand, serves to masquerade as damson, strawberry, raspberry, current, grape, etc., differing from each other only in color, but not as a rule in flavor. A variety of artificial colors are employed, mostly coal-tar dyes. To compensate for the lack of sweetness of the glucose, a minute quantity of one of the concentrated sweeteners, such as saccharin or dulcin, is sometimes added. Besides artificial colors, antiseptic substances are occasionally used, especially salicylic and benzoic acids.

All grades of apple stock are found in these preparations. A large source of supply is furnished by the parings and cores of canning establishments, to say nothing of the refuse of these factories, such materials being boiled with water, and the extract, variously colored to imitate the different fruits, being evaporated with commercial glucose.

Adulterated Jelly.—While it is easy to make an excellent apple jelly by simple evaporation of the pure apple juice, even without the addition of sugar, it is impossible, or at least difficult, to obtain the proper degree of stiffness with a mixture of apple stock and commercial glucose. It is customary, in the manufacture of cheap jellies, therefore, to employ what is technically termed a "coagulator." Formerly sulphuric acid, sometimes with addition of alum was used, but at present phosphoric acid is preferred. Citric or tartaric acid is also used for this purpose, as well as to increase the acidity. About 1% of the acid will cause the mass to gelatinize satisfactorily.

The lowest grade of apple jelly is made from the exhausted pomace, left as a residue after pressing out the juice for cider. Such stock is commonly mixed with water, and boiled down with glucose. Having been exhausted of its malic acid, pectose, and other soluble constituents, it lacks much of the flavor inherent in pure apple jelly. Various foreign gelatinizing agents are found in cheap jellies and preserves, such as starch, gelatin, and agar-agar. In the low-priced goods, starch paste has been employed. It should be remembered that starch exists in unripe apples, but hardly at all in the mature fruit, so that while mere traces of starch in jelly may be due to the use of green apples, its presence in large amounts is undoubted evidence of the admixture of starch paste.

Adulterated Jams.—Most of the cheap jams and bottled preserves sold on the market, though reinforced with apple stock, do in reality contain masses of fruit and berries of the kind stipulated on the label, as even a casual megascopic examination will show. That such low-priced preparations really contain genuine fruit pulp is not to be wondered at, when it is considered that much of the virtue of this fruit has sometimes been previously extracted by boiling, to produce fruit juices for higher-priced goods. Or, as in the case of jams containing strawberries, rasp-berries, and other small fruits with seeds, the juice is apt to have been previously expressed for pure jellies, while the residues are afterwards worked up with apple stock for low-priced jams. Hence the presence of pure fruit stock, or genuine berry seeds and pulp in jams, is in itself no criterion of purity, and, furthermore, it is unnecessary to use hay seed and other alleged foreign seeds as adulterants of cheap jam.

Compound Goods.—Many states have a law legalizing the sale of "compound" goods, providing they are distinctly so labeled. In other states, as, for instance, Massachusetts, the label must plainly state the name and percentage of the ingredients. In either case the analyst must discriminate, in classifying the inferior or low-grade preparations, between those that are labeled in accordance with the law, and those that are not. Only those not properly labeled can in such cases be classed as adulterated within the meaning of the law. Where such a law prevails, probably no class of food-products is so extensively affected by it as the low-grade jams, preserves, and jellies.

The restrictions as to labeling do not in all cases eliminate the element of deception. It is hardly justifiable, for example, to boldly label an alleged "currant jelly" which contains no currant, in the following manner:

Fruit juice	25%
Cane sugar	14%
Corn syrup.	61%
	~
	100%

The use of the term "fruit juice" surely implies to the unsuspecting purchaser that so much pure currant juice has entered into the jelly, elsewhere labeled in large letters "Currant," whereas all the juice is apple, and no currant juice has been used.

The following label is a type of those which discriminate between pure fruit and apple juice:

Fruit.	30%
Corn syrup	
Granulated sugar	
Apple juice	20%
	100%

Composition of Cheaper Grades.—Out of 66 samples of jellies, jams, and preserves analyzed by Winton, Langley, and Ogden in Connecticut, the samples being purchased in that state,* 17 samples contained starch paste, 35 were artificially colored with coal-tar dyes, and 19 contained salicylic or benzoic acid.

The following table has been compiled, showing the sugar content of some of the typical commercial jellies and jams analyzed in the laboratory of the Massachusetts State Board of Health. Nearly all of these were artificially colored, and found to contain little if any fruit, other than apple.

	Direct Polariza-	Invert Po	larization.	Per Cent	Per Cent Commer-
	tion.	At 20° C.	At 87° C.	Sucrose.	cial Glucose.
JELLY.	i				
Apple	+64.0	+ 28.0	+ 36.0	26.8	22. I
Currant A	+29.2	+20.0	+36.4	6.9	22.3
" B		+33.9	+40.8	5.7	25.0
Grape		+34.4	+46.0	20.6	28.2
Peach.		+108.8	+110.0	8.2	67.4
Pineapple	+114.0	+107.6	+110.0	4-9	67.4
Raspberry	+112.0	+92.0	+93.6	14.9	57-4
JAW.					
Damson A	+107.0	+94.4	+58.1	9-3	35.6
" B		+90.9	+83.6	3.2	51.2
Apricot	+99.0	+93.5	+85.6	4.1	52-4
Quince	+49.6	+43.6	+42.0	4.5	25.7
Raspberry A	+123.6	+119.2	+102.5	2.6	62.8
" <u>B</u>		+65.1	+46.9	9-3	28.7
''		+29.5	+37.2	27.2	22.8
Pineapple		+108.8	+110.0	8.2	67.4
Strawberry A		+21.3	+32.6	15.4	20.0
" B	+83.6	+72.0	+78.8	8.7	48.3

METHODS OF ANALYSIS.

As in the case of canned goods, but little information is to be derived as to adulteration of jams, jellies, and preserves by the ordinary determinations of moisture, ash, and nitrogen, and these are rarely made by the public analyst.

Of considerable importance in this regard, however, are the sugar determinations, made with a view to ascertaining the varieties of sugar

^{*} An. Rep. Conn. Exp. Sta., 1901, p. 130.

employed, as well as their approximate proportion in the products examined.

Total Solids.—Ten grams of the jam, which has been evenly pulped in a mortar, or 5 grams of the jelly, are weighed into a tared platinum dish, taking care to spread the sample as thinly as possible over the bottom of the dish, and dried to nearly a constant weight at 100°. Results yielded by this method, while sufficiently close for ordinary work, are not exact, due to the slight dehydration of the sugars. If extreme accuracy is required, dry *in vacuo* at 75° C., or in a McGill oven, page 586.

Soluble and Insoluble Solids.—A weighed amount of the evenly pulped sample, say 25 grams, is vigorously shaken in a 500-cc. graduated flask with water, preferably with the aid of a mechanical shaker, and, after filling to the mark, is again shaken. The residue is allowed to settle, and the supernatant liquid is decanted through a filter, and an aliquot portion of the filtrate, say 50 cc., is measured into a tared dish and evaporated to dryness, dried at 100° to a constant weight, and weighed for soluble solids. Insoluble solids are calculated by difference.

Ash.—The residue from the total solids is burnt at dull redness to an ash, cooled in a desiccator, and weighed.

Nitrogen is determined by the Gunning or Kjeldahl method, page 69, in from 5 to 10 grams of the uniformly mixed sample.

Determination of Sugars.—In products of the highest grade, wherein only cane sugar is employed, a large portion of the cane sugar is inverted in the process of boiling the jam or jelly, so that when the analyst examines it, he finds, as a rule, only a small amount of sucrose, and considerable invert sugar. It is possible, however, to calculate the amount of cane sugar originally employed, if such information is desirable. It is further of interest to calculate, at least approximately, the percentage of commercial glucose, when present, especially in cases where the package contains a formula setting forth the amount of the various ingredients used. In such cases the analyst is naturally called upon to verify the formula, since a wide variation in percentage composition from the statement on the label would constitute an offense under some state laws.

Polarization.—Use half the normal weight of the preserve or jelly for the Schmidt and Haensch instrument, viz., 13.024 grams in 100 cc. If fresh fruit or fruit juice is to be examined, use the full normal weight, 26.048 grams. Clarify, before making up to the mark, with subacetate of lead and alumina cream (using 2 to 3 cc. of each clarifier), filter, and obtain the direct reading; then invert in the usual manner, and obtain the invert readings at 20° C., and in the water-jacketed tube at 87° C., proceeding in detail as directed under honey, p. 641.

Calculation of Sugars.—Sucrose is determined by using Clerget's formula:

$$S = \frac{(a-b)_{100}}{_{142.66} - \frac{t}{2}}, \qquad (1)$$

This represents the sucrose actually present as such in the preserve or jelly, and not the amount originally used. If the latter is desired, it may be calculated from the formula,

$$S' = \frac{100b}{42.66 - \frac{t}{2}}, \qquad (2)$$

where S' is the per cent of cane sugar originally used, and b is the invert reading at f' of the normal solution.

If, after inversion, the correct reading at 20° is found to be 12 or more to the left of the zero, it can be safely inferred that no appreciable amount of commercial glucose is present, and it is unnecessary to make a third reading at 87°, unless to confirm the fact. In such a case, with cane sugar alone present, the reading at 87° will not, of course, vary much from o.

Invert Sugar.—In the absence of commercial glucose, the invert sugar is calculated as follows:

Invert sugar =
$$\frac{\text{(Sucrose-direct reading)}_{105.3}}{42.66 - \frac{t}{2}}, \dots (3)$$

or it may be determined directly from the copper reducing power.

Any decided reading above zero at 87° is due to the presence of commercial glucose, and when the latter is present, it is impossible to determine the invert sugar from the copper reduction or by formula No. 3. The following formula is proposed for calculating approximately the invert sugar from the polarization, in the presence of commercial glucose. While theoretically correct, the method is subject to practical limitations, which admit of only roughly approximate results in such mixtures as jelly or jam. It is perfectly accurate only in mixtures of sucrose, glucose, and invert sugar.

Invert sugar =
$$\frac{\left(\frac{\text{Reading due to glucose and}}{\text{inverted sucrose at } P}\right) - \left(\frac{\text{Invert reading}}{\text{at } P}\right)}{\pm \left(42.66 - \frac{t}{2}\right)}$$
 105.3 (4)

These formulas, (3) and (4), serve at best to indicate the approximate amount of invert sugar present in the sample, resulting from the inversion of a portion of the original sucrose in the natural process of manufacture of the jam or jelly, and not the total invert sugar resulting from the inversion by the analyst of all the sucrose.

The factor 105.3 is used, since, in the natural process of inversion, 100 parts of sucrose become 105.3 parts of invert sugar.

Example.—The invert sugar in the sample of apple jelly first on the list in the table on page 916 is calculated as follows:

Invert reading at $f''(20^\circ) = 28.0$. Reading due to glucose at $20^\circ = .221 \times 175 = 38.68$. " " inverted sucrose at $20^\circ = .268 \times -34 = -9.11$. Invert sugar = $\frac{(38.68 - 9.11) - 28}{28,66}$ 105.3 = 5.76%.

Reducing Sugar by Copper Reduction.*—Five grams of the preserve or jelly (or 25 grams of the fresh fruit or fruit juice) are transferred to a 100 cc. graduated flask, clarified by the addition of 2 or 3 cc. each of subacetate of lead and alumina cream, made up to the mark, shaken, and filtered. An aliquot part of the filtrate is then measured into another 100 cc. flask, and treated with enough of a saturated solution of sodium sulphate to precipitate the lead, after which it is made up to the mark and filtered. The amount of sugar solution measured off into the second flask is such that, when finally made up to 100 cc. as described, approximately 1 of 1% of reducing sugar is present, as roughly estimated by the total solids and polarizations. The reducing sugar is then determined in the filtrate as dextrose by Defren's method, page 594, or if Allihn's method is used (p. 608) the amount of reducing sugar present should approximate 1%.

Commercial Glucose.—While it is impossible to determine the exact percentage of this substance in preserves and jellies, by reason of the varying composition of its component parts, it is quite feasible to approximate very closely to the amount present. Indeed, this approximate

^{*} U. S. Dept. of Agric., Bur. of Chem., Bul. 65, p. 78.

method of calculation, wherein glucose is treated as a chemical entity, has been found in practice to be much more close to the actual truth than results gained by methods wherein the copper reducing power enters as a factor, or methods for determining separately dextrin, maltose, and dextrose. Calculate the commercial glucose in jellies and jams exactly as in the case of honey, p. 642.

Dextrin.*—If alcohol be added to a somewhat thick solution of the fruit product, a white turbidity is at once apparent, followed by the formation of a thick gummy precipitate, if dextrin is present. In the absence of dextrin there is no turbidity, but a light flocculent precipitate.

To determine the dextrin, dissolve † 10 grams of the sample in a 100-cc. flask; add 20 mg. of potassium fluoride, and then about one-quarter of a cake of compressed yeast. Allow the fermentation to proceed below 25° C. for two or three hours to prevent excessive foaming, and then place in an incubator at a temperature of from 27° to 30° C. for five days. At the end of that time clarify with lead subacetate and alumina cream; make up to 100 cc. and polarize in a 200-mm. tube. A pure fruit jelly will show a rotation of not more than a few tenths of a degree either to the right or to the left. If a Schmidt and Haensch polariscope be used, and a 10% solution be polarized in a 200-mm. tube, the number of degrees read on the sugar scale of the instrument, multiplied by 0.8755, will give the percentage of dextrin, or the following formula may be used:

Percentage of dextrin=
$$\frac{C \times 1000 \times V}{198 \times L \times W}$$
,

in which

C = degrees of circular rotation,

V = volume in cubic centimeters of solution polarized,

L = length of tube in centimeters,

W = weight of sample in solution in grams.

Determination of Tartaric Acid.‡—To 100 cc. of the fruit juice add 2 cc. of glacial acetic acid, 2 or 3 drops of a 20% potassium acetate solution, and 15 grams of pure finely powdered potassium chloride; dissolve this by shaking, and then add 20 cc. of 96% alcohol. Then stir vigorously for one minute, rubbing the walls of the

^{*} Bur. of Chem., Bul. 65, p. 78.

[†] Bigelow and McElroy, Jour. Am. Chem. Soc., 1893, 15, 668.

[‡] Halenke & Moslinger, Zeit. anal. Chem., 1895, 34, 283; Bur. of Chem., Bul. 65, p. 80.

beaker with the glass stirring-rod to start the crystallization of the potassium bitartrate. Allow to stand fifteen hours at room temperature. Filter, and wash the precipitate into a Gooch crucible with a thin asbestos felt, using the vacuum pump. Wash with a mixture of 15 grams of potassium chloride, 20 cc. of alcohol, and 100 cc. of water. The beaker is rinsed three times with a few cubic centimeters of this solution. The precipitate is also washed with a few cubic centimeters, but so that not more than 20 cc. in all of the wash solution is used. The precipitate and asbestos filter are washed back into the beaker, and heated to boiling. While still hot, the solution is titrated with decinormal alkali, using phenolphthalein as indicator. To the amount of alkali used must be added 15 cc. for the potassium bitartrate remaining dissolved in the solution. 1 cc. of decinormal alkali is equivalent to 0.0150 grams of tartaric acid.

Determination of Citric Acid.*—Fifty cubic centimeters of the fruit solution is evaporated on the water-bath to a syrupy condition. To the residue add, very slowly at first, stirring constantly, 95% alcohol until no further precipitate is formed; 70 to 80 cc. are generally enough. Filter, and wash the residue with 95% alcohol. Evaporate the filtrate to eliminate the alcohol, take up the residue with a little water, and transfer to a graduated cylinder, making up to 10 cc. To 5 cc. of this solution, add half a cubic centimeter of glacial acetic acid, and to this add, drop by drop, a saturated solution of lead acetate. The presence of citric acid is shown by the appearance of a precipitate, which possesses the property of disappearing on being heated, and reappearing on cooling. In order to separate the citric acid from other acids, heat to boiling, filter, and wash with boiling water; then allow to cool, and the precipitate of lead citrate will re-form. This lead precipitate may be filtered off, washed with weak alcohol, dried, weighed, and the citric acid calculated. It is necessary that there shall be no tartaric acid present. If the tartaric acid has been estimated, any error on this account may be avoided by adding enough decinormal potash to neutralize the tartaric acid before the alcohol is added.

Detection of Coloring Matter.—Boil white woolen cloth or worsted in a solution of the jelly or jam, acidified with hydrochloric acid, or with acid sulphate of potassium, according to Arata's method and test for the color on the dyed fabric by methods given in detail in Chapter XVII.

^{*} Moslinger, Zeit. Unter. Nahr. u. Genuss., 1899, 2, p. 93; U. S. Dept. of Agric., Bur. of Chem., Bul. 65, p. 80.

Detection of Preservatives and Concentrated Sweeteners.—Extract an acid aqueous solution of the fruit product with ether or chloroform in a separatory funnel, and test for benzoic and salicylic acids and for saccharin in the ether extract. If dulcin is suspected, extract with acetic ether.

Detection of Starch.*—Heat an aqueous solution of the preserve or jelly nearly to the boiling point, and decolorize by the addition of several cubic centimeters of dilute sulphuric acid and afterwards permanganate of potassium. This treatment does not affect the starch, which is tested for with iodine in the ordinary manner in the solution after cooling. In the clear filtrate from a boiled apple pulp solution, free from added starch, little or no darkening should occur on the addition of the iodine reagent. If, however, the reagent is added to the residue of the previously boiled pulp, the presence of starch inherent in the apple is usually recognized by the blue color produced thereon.

The presence of any considerable added starch paste in a fruit preparation is thus readily indicated by an intense blue color obtained by adding the iodine reagent to the filtrate (free from fruit pulp).

Detection of Gelatin.—Robin's Method.†—Add to a thick aqueous solution of the preserve or jelly sufficient strong alcohol to precipitate the gelatin. Decant the supernatant liquid after settling, set aside part of the precipitate, and dissolve the remainder in water. Divide the latter solution in two parts, to one of which add, drop by drop, a fresh solution of tannin, which precipitates gelatin if present. To the remainder add picric acid solution, which in presence of gelatin forms a yellow precipitate. The portion of the yellow precipitate set aside is transferred to a test tube, and heated over the flame with a little quicklime. If gelatin is present, ammonia will be given off, apparent by the odor, and by fumes of ammonium chloride when a drop of hydrochloric acid on a glass rod is held at the mouth of the bottle.

Leffmann and Beam's Method. ‡—Boil the sample with water, filter, and boil the filtrate with an excess of potassium bichromate. Cool, and add a few drops of sulphuric acid. A flocculent precipitate indicates gelatin.

Detection of Agar Agar. \[
\]—The jelly is heated with 5% sulphuric acid, a little potassium permanganate is added, and, after settling, the

^{*} U. S. Dept. of Agric., Bur. of Chem., Bul. 65, p. 81.

[†] Girard et Dupré, Analyse des Matières Alimentaires, p. 578.

[‡] Select Methods of Food Analysis, p. 324.

[§] Marpmann, Zeit. f. angew. Mikrosk, 1896, p. 260; U. S. Dept. of Agric., Bur. of Chem., Bul. 65, p. 81.

sediment is examined by the microscope for diatoms, which will be found in large numbers if agar agar has been used.

Detection of Apple Pulp.—A distinct clue to the presence of apple pulp in fruit preparations is often furnished by the characteristic apple odor given off when a small amount of the sample is heated to boiling with water in a test tube. Under such conditions, the apple odor is quite apparent, as distinguished from that of other fruits, especially if the apple is the chief fruit present, or predominates in the mixture.

Apple pulp in fruit preserves, free from added starch, may usually be recognized by a microscopical examination, using iodine reagent. The cell contents of the pulp will show the characteristic blue color, undoubtedly due to portions of unconverted starch still remaining in them.

Detection of Fruit Tissues under the Microscope.—It is a matter of some difficulty, by means of a microscopical examination, to identify with certainty the various fruits and vegetables that might be used in a jam or jelly as adulterants. The structural features of the common fruits, while possessing distinctive points of difference when examined separately and in the raw products, are so changed or broken down by the process of cooking, as to be with difficulty recognizable. The soft parenchyma which forms the main portion of the tissue of the fruit pulp is, as a rule, more or less disintegrated.

In the case of some of the smaller fruits, as the currant and raspberry, the cuticles resist the cooking process to such an extent as to show characteristic fragments, often recognizable in preserves and jellies under the microscope. The seeds are highly characteristic.

FRUIT JUICES.

Sweet cider, orange juice, lime juice, grape juice, raspberry shrub, and the juices of various other fruits and berries, may be so prepared and sterilized as to keep without fermentation when bottled, and are so put up in considerable variety, either with or without the addition of sugar.

Such preparations, if of the highest purity, should consist of the undiluted juices of these fruits, separated by pressure and carefully sterilized and bottled. They should contain no other fruit juice than that specified on their labels, and should be free from alcohol, added antiseptics, or coloring matter, unless the label specifies the presence of the added foreign materials. The addition of pure cane sugar to such prepara-

tions as grape juice is allowable, as well as charging with carbon dioxide to form so-called carbonated drinks.

The following analyses of pure fruit juices are taken from tables prepared by Winton, Ogden, and Mitchell, showing results on samples purchased in the Connecticut market, as well as on some samples made in the laboratory.*

5	Solids.	Acids Other					A CARLL	zation.	
	onus.	than CO ₂ as Citric.	Cane Sugar.	Invert Sugar.	Direct.	After Inver- sion.	Temper- ature C.	Invert Reading at 86° C.	
COMMERCIAL PRUIT									
JUICES.							!		
Blackberry	5.32	0.65	0.0	4.6	-1.3	-1.3	29.0		
	14.33	0.80	0.0	6.5	-1.9	-1.9	26.0		
	10.00	2.41	0.0	9.2	-2.7	-2.7	26.0	,	
Red currant	7.58	2.09	0.0	7.2	-2.1	-2.1	27.0		
	15.29	0.91	0.0	21.7	-6.5	-6.5	25.0	-1.0	
Lime fruit	7.78	6.50	0.0	0.0	0.0	0.0			
	12.72	2.44	0.0	7.1	-2.1	-2.1	26.0	}	
Pineapple	8.07	0.81	1.5	5.1	0.0	-2.0	26.0		
	10.81	1.00	0.0	0.3	-0.1	-0.1	26.0		
	10.41	0.99	0.0	16.7	-5.0	-5.0	25.0	-2.2	
Black raspberry	8.47	1.36	0.0	7.8	-2.3	-2.3	26.0		
Strawberry	5.69	0.99	0.0	5.1	-1.5	-1.5	26.0		
MADE IN LABORA- TORY.									
Peach.	12.70	0.95	5-4	2.1	4.8	-2.2	28.0	0.0	
Red raspberry	9.41	1.19	ŏ.8	8.6	-i.6	-2.8	26.0	0.0	
Blackberry	8.94	1.22	0.0	8.7	-2.4	-2.4	30.0	0.0	
	11.40	0.51	0.6	16.7	-4.0	-4.8	30.0	-1.0	
- · · · ·	13.90	o.ĕ8	7-4	9.1	4.7	-4.8	28.0	-o.8	

Antiseptics found most commonly in these preparations are boric, salicylic, benzoic, and sulphurous acids. Beta-naphthol should also be looked for. For methods of separation and examination see Chapter XVIII.

Unfermented Grape Juice has the following average composition: †

	Austria, Per Cent.	California, Per Cent.
Solid contents by spindle (Balling). Alcohol Total acid (as tartaric). Volatile acid. Grape sugar. Cream of tartar. Free tartaric acid. Ash. Phosphoric acid.	21.62 None .78 .01 19.62 .61 .03 .37	20.60 None -53 .03 19.15 -59 .07 .19

^{*} An. Rep. Conn. Exp. Sta., 1899, p. 136.

[†] California Exp. Sta., Bul. 130.

Grape juice is prepared by sterilizing at a temperature of 80° the juice expressed from the crushed grapes, filtering by means of a press or otherwise, and sealing in carefully sterilized bottles. After bottling, a final sterilization is conducted at a temperature 5° below the first. Bottled grape juices are rarely carbonated.

Bottled Sweet Cider.—The composition of pure, freshly expressed apple juice is shown by the following table of analyses by Browne:*

	Specific Gravity.	Solids.	Invert Sugar.	Su- crose.	Total Sugar.	Total Sugar after Inver- sion.	Free Malic Acid.	Ash.	Unde- ter- mined (Pectin, etc.).	Left- handed Rotation Degrees Ventake 400 mm. Tube.
Red astrachan										23.72
Early harvest Yellow transparent.										24.32
Sweet bough					10.14				0.44	39.40
Baldwin, green										36.16
"ripe Ben Davis										49.00

Bottled sweet cider, properly sterilized, should not differ materially from the fresh juice, and should contain no alcohol.

Salicylic acid is the antiseptic most commonly found in sweet bottled ciders examined by the writer.

Lime or Lemon Juice.—This, according to the U. S. Pharmacopœia, should consist of the freshly expressed juice of the ripe fruit of Citrus limonum (Risso), natural order of Rutaceæ. Our supply of both lemons and limes comes chiefly from the West Indies and the Mediterranean. Both varieties of the genus Citrus are used indiscriminately for furnishing commercial lime juice, though strictly speaking, only that of the lemon is recognized in the Pharmacopœia. The juice is sharply acid, and is largely composed of citric acid (about 7%), gum, sugar (3 to 4 per cent), and inorganic salts from 2 to 2½ per cent. It also usually contains a little lemon oil from the rind. According to the pharmacopœia, lemon juice (Limonis succus) should conform to the following requirements:

"It has an acid reaction upon litmus paper, due to the presence of about 7% of citric acid.

[&]quot;Specific gravity: not less than 1.030 at 15° C.

^{*} Penn. Dept. Agric., Bul. 58, p. 29.

"On evaporating 100 grams of the juice to dryness, and igniting the residue, not more than 0.5 gram of ash should remain."

Of thirty samples of commercial lime juice examined in the Massachusetts State Board of Health laboratory, representing fifteen brands, all were deficient in citric acid, containing from 1.92 to 4.15 per cent, thus showing that these preparations are frequently watered. Fifteen were found to contain salicylic acid, seven had sulphurous acid, while two contained both these preservatives. Several were found colored with coal-tar dyes.

One sample examined by the author, purporting to be a "pure West Indian Lime Juice, triple refined," proved to be a mixture of hydrochloric and salicylic acids, colored with a coal-tar dye, and contained no lime juice whatever.

Acidity of lime juice is obtained by titrating 6.8 cc. of the sample against tenth-normal sodium hydroxide with phenolphthalein. The number of cubic centimeters of the standard alkali required, divided by 10, gives the per cent of citric acid present.

FRUIT SYRUPS.

Two classes of these preparations are on the market, one for use in soda-fountains, and one for "family trade," intended as a basis for sweetened drinks to be diluted with water and sugar. Some are made exclusively from pure fruit pulp and sugar, sterilized by heating and put up in tightly sealed bottles, while others of the cheaper variety are more apt to be entirely artificial both in color and in flavor, deriving the latter principally from the wide variety of artificial fruit essences now available. Commercial glucose is a frequent ingredient. The same classes of coaltar dyes and antiseptics are found in these preparations as in the other fruit products. Fruit syrups are frequently found to contain such materials as gum arabic and quillaia, or soapbark, used both for a thickener, and to give a "bead" or froth when used in soda water, and in connection with carbonated drinks.

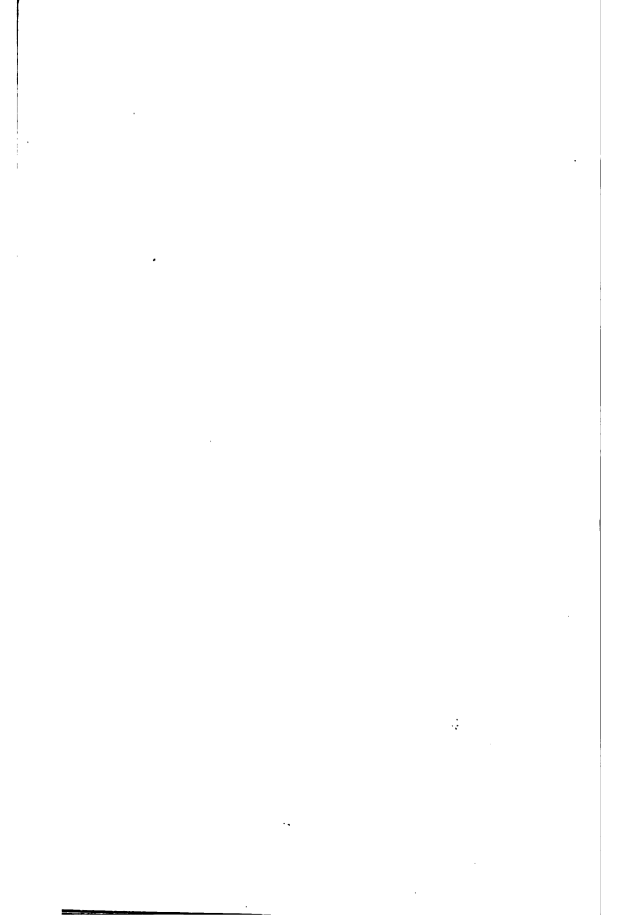
For purposes of comparison with such fruit-pulp preparations as may come to the analyst for examination, he is referred to the analysis of fruits found on page 274.

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INDEX.

Abbé refractometer, 100, 108 construction, 109	Alcoholic beverages, 653, 654. See also Liquors.
influence of temperature, 110	references on, 756
manipulation, 109	state control of, 654
	toxic effect of, 655
Abrastol, 837	
Absinthe, 754	fermentation, 653
Acetanilide in vanilla extract, 858	Aldehydes, determination, 745
tests for, 859	Ale, 709, 712. See also Beer.
Acetyl value, 497	Aleurone, 90
Achroodextrine, 575	Alkaloidal nitrogen, 40, 46
Acid fuchsin, 797	Alkaloids, proof of absence of, 726
Acids, fatty, 481, 484, 499, 500	Alkanna tincture, 92
of acetic series, 471	Allantoin, 299
of linoleic series, 472	Allen-Marquardt method for fusel oil, 747
of oleic series, 472	Allihn's sugar method, 608
mineral, in vinegar, 766, 767	tables, 609
organic, 47	Allspice, 420
Ackermann and Steinmann's table for	adulteration, 424
alcohol from refraction, 715	composition of, 420
Ackermann's table for extract from refrac-	microscopical structure, 422
tion, 721	standard, 424
Adams' fat method, 134	tannin in, 421
"Aerated" butter, 540	Almond extract, 873
Agar agar, in jelly, 914, 922	adulteration of, 875
Aging of liquors, 731, 732	alcohol in, 877
Albumin, acid, 44	benzaldehyde in, 874, 875
alkali, 44	hydrocyanic acid in, 874,
determination in milk, 146	877
of muscle, 211	nitro benzol in, 876
preparation of, 263	standards, 874
Albuminoids, 42	meal, 358
Albumins, 41, 207	Almonds, bitter, oil of, 873, 874
Albumose, 44, 45	Alum in baking powder, 333, 344
Alcohol, detection, 657	in bread, 326
determination, 658	in flour, 315
by distillation, 658	in pickles, 910
by ebulioscope, 675	Alumina, determination of, 344
by evaporation, 660	Aluminum salts in balina namen
	Aluminum salts in baking powder, 344
from refraction, 715	in cream of tartar, 344
from specific gravity, 658, 659	Amagat and Jean's refractometer, 100
in malt liquors, 715	Amides, 45
extract of spices, 470	in milk, 147
methyl-, 749, 869	Amido nitrogen determination, 74, 147
preparation of, 730	in wheat, 299
stills, 659	Amino acids, 40, 45
tables, 661-674	Ammonia, determination, 74
•	, 93 1

932 INDEX.

	•
Ammonia, in baking powder, 346	Balances, 20
in foods, 40, 46	Bamihl test for gluten, 322
in milk, 147	Banana essence, artificial, 884, 885
Ammonium fluoride, 835	Barium compounds in colors, 782
Amthor test for caramel, 752	Bark as an adulterant, 428
Amylodextrin, 575	Barley, 271, 272
Amyloid, 91, 92	ash, 302
Analyst, functions of, 3, 4	microscopy of, 309
Angostura, 754	proteins, 300
Anilin orange in milk, 177	starch, 281
Animal diastase, 284	Basic colors, 793, 796
Anise extract, standards, 880	Baudouin's sesame oil test, 519
oil, standards, 880	Beading oil, 738
Annatto in butter, 536, 537	Beans, 272, 388
in milk, 175, 177	Beaumé and Brix scales compared, 617-620
tests for, 789	Bechi's cottonseed oil test, 517
Antiseptics, see preservatives	Beckman's test for glucose in honey, 641
Apparatus, 20	Beef, composition of, 213
Apple essence, imitation, 884, 885	cuts of, 213
Apple juice, 680	stearin, microscopical structure, 558
Apples, composition of, 274, 275	tallow 529
Araban, 285, 288	Beer, 707
Arabinose, 285, 288	acids in, 724
Arata's color test, 794	adulteration of, 711
Army rations, 257	alcohol in, 715
Arsenic detection and determination, 75, 76	aloes in, 727
compounds in colors, 783	arsenic in, 713, 728
Gutzeit test for, 632	ash of, 714
in beer, 713, 728	bitter principles of, 726
in vinegar, 778	bock-, 709
Marsh apparatus, 75	brewing of, 708
Artificial colors, 780	carbon dioxide in, 726
fruit essences, 884, 885	chiretta in, 711, 727
sweeteners, 842	composition of, 700
references on, 847	degree of fermentation of, 724
	dextrin in, 724
Asherton Shor preparation of ros ros	extract gravity of, 722
Asbestos fiber, preparation of, 594, 598 Ash analysis, scheme for, 301	extract in, 715
determination of, 61	specific gravity method, 722
of food, 47	refractometer method, 722
· '•	gentian bitter in, 711, 727
Asparagin, 45, 299	glucose in, 710
Auramin, 782	
Babcock asbestos milk fat method, 135	glycerin in, 724
	lager-, 708
milk solids method, 134	methods of analysis, 714
centrifugal fat method, 136	phosphoric acid in, 725
milk formulæ, 153	preservatives in, 713, 729
test bottles, 138	proteins of, 725
Baier and Neuman's test for sucrose in milk,	quassiin in, 711, 727
197 Paking pourders and	references on, 756
Baking powders, 332	schenk-, 708
adulteration of, 334	standards, 711
alum, 333, 334	temperance-, 714
methods of analysis, 336	varieties of, 708
phosphate, 334	uno-, 714
tartrate, 332	weiss-, 709

Day much not	Donal along to and
Beer, wort, 708	Bread, alum in, 326
gravity of, 722	baking of, 323
Beeswax, 643	composition of, 324, 325
refractometer reading of, 645	fat in, 326
Beet (color), 788	Breakfast cereals, 352
sugar, 569	Brewing beer, 708
Bellier's peanut oil test, 524	Brie cheese, 202
Benches, 15	Brix scale compared with Beaumé, 617-620
Bénédictine, 754	Bromination oil test, 494
Benzaldehyde, 874, 875	Bromine absorption of oils, 492
artificial, 874	Brown and Duvel's method for moisture in
in almond extract, 875	grain, 278
Benzoic acid, 827	Brown colors, 784, 786, 808
detection of, 828	sugar, 568
determination, 830	Browne's method for dextrin in honey, 640
in milk, 180	test for invert sugar in honey, 642
Betaine, 45, 299	Brucke's glycogen method, 236
Beta-naphthol, 837	
	reagent, 236
Bigelow and McElroy's cane-sugar method,	Buckwheat, 271
192 Tells (-1 -)00	ash of, 302
Bilberry (color), 788	composition of, 271, 272
Birotation, 584, 639	flour, 313
Biscuit, gluten, 358	microscopy of, 311
soja bean, 358	Burgundy wine, artificial, 692
Bisulphites as preservatives, 833	Butter, 201, 529
Bitter almonds, oil of, 873	adulteration of, 535
Biuret reaction, 41	annatto in, 536
Blackberry (color), 788	ash in, 534
Blarez test for fluorides, 835	azo colors in, 536, 537
Blast pump, 19	boric acid in, 538
Blue colors, 783, 784, 786, 792, 810	carrotin in, 536
"Blown" cans, 890	casein in, 534, 551
Bock-beer, 709	coloring in, 535
"Boiled" butter, 540	composition of, 530
Bombay mace, 467	distinction from oleomargarine and
Bomb calorimeter, 47	process butter, 546
Bömer's phytosterol acetate test, 507	effects of feeding, 531
Borax, 821	
	fat, composition of, 530
Boric acid, 821	standard, 535
detection, 182, 184, 822	fat in, 533, 534
determination, 821, 823	filled, 540
in butter, 538	foam test, 549
in meat, 220,232	formaldehyde in, 539
in milk, 182, 184	glucose in, 539
Bourbon whiskey, 732, 734, 737	methods of analysis, 531
Brandy, 739	microscopical examination of, 552
adulteration of, 741	milk test, 550
composition of, 739	preservatives in, 538
"drops," 649	references on, 562
methods of analysis, 745	renovated, 540
new, 740	salicylic acid in, 539
potable, 740	salt in, 534
standards, 740	standard, 535
Bread, 317, 323	sulphurous acid in, 539
acidity of, 325	water in, 531
adulteration of, 326	turmeric in, 536
manterpriori ori Dec	

Butter, Waterhouse test, 550	Canned food, impurities in, 890
Butterine, 541	metallic impurities in, 892
Butterine oil, 522	method of canning, 888
Butyro-refractometer, 100, 101	methods of analysis, 889
critical line of, 106	references on, 927
limits of butter readings, 547	Canned fruits, 887, 889
manipulation, 102	meats, 22
oil readings on, 478, 479	vegetables, 887, 889
olive and cottonseed oil readings, 514	Cans, detection of spoiled, 890
sliding scale for, 107	gases from spoiled, 891
special thermometer for, 549 table of equivalent refractive indices,	Capers, 909
104, 105	Capsicin, 440
temperature variation of reading,	Capsicums, 439 Caramel, 790
107	in distilled liquors, 752
testing scale, 104	in milk, 176, 177
testing scare, 104	in vanilla extract, 860
Caffeine, 372	in vinegar, 777
determination of, 373, 384	Carbohydrates, 46, 47, 74, 279
in cdcoa, 400	of cereals, 279, 295
Caffeol, 379	of eggs, 263
Caffetannic acid, 379, 382	Carbon dioxide in baking chemicals, 336
Cake, 327	in beer, 726
Calcium carbonate crystals, 90	in yeast, 330
oxalate crystals, 90	Carnin, 211
sucrate, 196	Carrot (color), 789
California wines, 688	Casein, 43, 125, 126
Calorie, 47, 48	determination in milk, 145
Calorimeter, bomb, 47	Caseose, 44
oil, 495	in cheese, 203
respiration, 2	in milk, 146
Camembert cheese, 202	Casoid flour, 358
Camera, 96	Cassia, 424
Canada balsam, 86	adulteration of, 428
Candy, see confectionery	buds, 425
standard, 645	composition of, 422
Cane sugar, 566	extract, standards, 830
ash of, 567	microscopical structure, 426
composition of, 568	oil, 425, 880
detection of, 585	standards, 880
in milk, 197	standard, 428
determination of,	Cayenne, 439
by copper reduction, 590, 612	adulteration of, 443
by polarimetry, 586, 614	coal-tar colors in, 444
in cereals, 295	colors in, 444
inversion of, 588, 589	composition of, 441
manufacture of, 567	microscopical structure, 441
methods of analysis, 585	mineral adulterants in, 444
moisture in, 586	oil of, 440
quotient of purity, 586	redwood in, 444
refining, 570	standard, 443
test for, 585	Cazeneuve's color scheme, 705, 706
Canned food, 887	Celery seed extract, standards, 880
antiseptics in, 903	oil, standards, 880
composition of, 889	Cellulose, 47, 285
decomposition of, 890	Centrifuge, milk-fat, 136, 137

Centrifuge, universal, 25 Cereal products, microscopy of, 305 Cereals, 271	Cholesterol, crystallizations of, 504 determination of, 503 distinction from phytosterol,503
ash of, 302	separation of, 503
breakfast foods, 352	Cholin, 45, 299
cane sugar in, 295	Chromate of lead, 647,
carbohdyrates of, 279	Chromogenic bacteria, 130
separation of, 295	Cider, 678
composition of, 271	adulteraton of, 682
crude fiber in, 277, 296	ash of, 682
dextrin in, 295	composition of, 679
hemicelluloses in, 296	fermented, 680
methods of proximate analysis, 276	malic acid in, 683
pentosans in, 285, 296	manufacture of, 678
proteins of, 296	methods of, analysis, 606
references on, 361	references on, 757
starch determination in, 283, 296	sweet, 925
Chace's citral method, 866	
Champagne, 687	vinegar, 760, 771
Chaptalizing, 693	watering of, 682
	yeast in, 678
Chartreuse, 754	Cinnamon, 424
Cheddar cheese, 202	composition of, 425, 426
Cheese, 201	extract, 881
adulteration of, 203	microscopical structure of, 426
amides in, 206	oil, standards, 881
ammonia in, 206	standard, 428
ash in, 204	Citral, 871
composition of, 201	determination, 866
cream, 203	Citric acid in fruit products, 921
lactic acid in, 207	in lime juice, 925
fat in, 204, 207	in milk, 126, 127
filled, 203	Citronellal, 872
methods of analysis, 204	Citronella oil, 871, 872
milk sugar in, 207	Clams, 256
nitrogen compounds of, 205	Claret wine, 687
paranuclein in, 206	Clarifying reagents in microscopy, 92
peptones in, 206	in sugar analysis, 586,614
proteins in, 205	Clerget's formula, 588
sampling, 204	Clove extract, 881
skimmed milk, 203	oil, 881
standards, 203	Cloves, 412
varieties of, 202	adulteration of, 418
water in, 204	composition of, 414
whole milk, 203	cocoanut shells in, 419
Chicory, 386, 388, 389	exhausted, 418
Chili sauce, 907	microscopical structure, 416
Chilton cheese, 202	oil of, 881
Chiretta, 727	standard, 418
Chlor iodide of zinc, 91	stems, 417
Chloral hydrate, 93	tannin in, 415
Chlorine in vegetable substances, 305	Clupein, 43
Chocolate, see cocoa,	Coal-tar colors, 791
milk, 397	acid, 796
composition of, 397	allowed, 792
sucrose and lactose in, 399	Arata's test, 794
Cholesterol, 502	basic, 793

Coal-tar colors, classification, 791, 798, 800 detection of, 793 double dyeing method, 794 dyeing wool by, 793 extraction by amyl alcohol, 795 identification of, 793, 797, 799, 803 in milk, 177 in sausages, 239 Rota's scheme for, 797 Sostegni and Carpentieri's color test, 794 separation with ether, 796 Cochineal, 790 in sausages, 238 Cocoa, 392, 393 adulteration of, 402 alkali in, 403 ash of, 396 butter, 393, 529 caffeine in, 400 composition of, 393 foreign fat in, 402 manufacture of, 393 methods of, analysis, 398 microscopical structure, 403 nibs, 394, 402 nitrogeneous bodies in, 396 pentosans in, 396 references on, 406 shells, 394, 395, 495 standards, 402 starch in, 394, 395, 399, 495 sugar in, 399, 495 theobromine in, 396, 400 Cocoanut oil, 528 pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffeine in, 380, 384 coloring of, 384 composition of, 379 fat in, 579 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384 references on, 406		
detection of, 793 double dyeing method, 794 dyeing wool by, 793 extraction by amyl alcohol, 795 identification of, 793, 797, 799, 803 in milk, 177 in sausages, 239 Rota's scheme for, 797 Sostegni and Carpentieri's color test, 794 separation with ether, 796 Cochineal, 790 in sausages, 238 Cocoa, 392, 393 adulteration of, 402 alkali in, 403 ash of, 396 butter, 393, 529 caffeine in, 400 composition of, 393 foreign fat in, 402 manufacture of, 393 methods of, analysis, 398 microscopical structure, 403 nibs, 394, 402 nitrogeneous bodies in, 396 pentosans in, 396 pentosans in, 396 pentosans in, 396, 400 Cocoanut oil, 528 pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffeine in, 396, 385 caffeine in, 396, 385 coffeine, 379 adulteration of, 379 adulteration of, 384 caffeine in, 394, 395, 399, 405 sugar in, 399, 405 theobromine in, 396, 400 Cocoanut oil, 528 pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 caffeine in, 380, 384 composition of, 379 fat in, 379 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384 microscopical structure, 3	Coal-tar colors, classification, 791, 798, 800	Coffee, standards for, 384
dyeing wool by, 793 extraction by amyl alcohol, 795 identification of, 793, 797, 799, 803 in milk, 177 in sausages, 239 Rota's scheme for, 797 Sostegni and Carpentieri's color test, 794 separation with ether, 796 Cochineal, 790 in sausages, 238 Cocoa, 392, 393 adulteration of, 402 alkali in, 403 ash of, 396 butter, 393, 529 caffeine in, 400 composition of, 393 foreign fat in, 402 manufacture of, 393 methods of, analysis, 398 microscopical structure, 403 nibs, 394, 402 nitrogeneous bodies in, 396 pentosans in, 396 references on, 406 shells, 394, 395, 405 standards, 402 starch in, 394, 395, 405 standards, 402 starch in, 394, 395, 495 theobromine in, 396, 400 Cocoanut oil, 528 pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffein in, 380, 384 caffetannic acid in, 379, 382 caffein in, 389, 384 composition of, 379 fat in, 379 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384 microscopical structure, 386 "pe		
extraction by amyl alcohol, 795 identification of, 793, 797, 799, 803 in mills, 177 in sausages, 239 Rota's scheme for, 797 Sostegai and Carpentieri's color test, 794 separation with ether, 796 Cochineal, 790 in sausages, 238 Cocoa, 392, 393 adulteration of, 402 alkali in, 403 ash of, 396 butter, 393, 529 caffeine in, 400 composition of, 393 foreign fat in, 402 manufacture of, 393 methods of, analysis, 398 microscopical structure, 403 nibs, 394, 402 nitrogeneous bodies in, 396 pentosans in, 396 references on, 406 shells, 394, 395, 495 standards, 402 starch in, 394, 395, 399, 405 sugar in, 399, 405 sugar in, 399, 405 theobromine in, 396, 400 Cocoanut oil, 528 pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffeine in, 380, 384 caffetannic acid in, 379 chicory in, 388, 389 coloring of, 384 composition of, 797 Colors, artificial, 780 acid fuchsin 797 allowed, 792 animal, 790 arsenic compounds, 783 barium compounds, 783 barium compounds, 782 basic, 793 blue, 793, 784, 786, 792, 810 brown, 784, 786, 808 caramel, 790 coherent, 793 cochineer, Schreiner's, 77 Colors, artificial, 780 acid fuchsin 797 allowed, 792 animal, 790 arsenic compounds, 783 barium compounds, 783 barium compounds, 783 barium compounds, 782 basic, 793 blue, 793, 784, 786, 792, 810 brown, 784, 786, 808 caramel, 790 coherowric analysis, 77 Colors, artificial, 780 acid fuchsin 797 allowed, 792 animal, 790 coherowric analysis, 79 colormetric analysis, 77 Colors, artificial, 780 acid fuchsin 793 allowed, 792 animal, 790 coherowric, 783 barium compounds, 783 barium compounds, 783 barium compounds, 783 barium compounds, 782 basic, 793 blue, 793, 784, 786, 808 caramel, 790 coherometric analysis, 792 colorimetric analysis, 792 colorimetric analysis, 792 colorimetric, 204 compounds, 782 basic, 793 cochineal, 790 copression of 184, 402 introgeneous bodies in, 396 green, 783, 784, 785, 792, 803 indigo, 790, 810 injurious, 782, 784 identification of, 793, 797, 803 indigo, 790, 810 injurious, 782, 783 in butter, 335 in cayenne, 444 in co	double dyeing method, 794	substitutes, 392
identification of, 793, 797, 799, 803 in milk, 177 in sausages, 239 Rota's scheme for, 797 Sostegni and Carpentieri's color test, 794 separation with ether, 796 in sausages, 238 Cocoa, 392, 393 adulteration of, 402 alkali in, 403 ash of, 396 butter, 393, 529 caffeine in, 400 composition of, 393 foreign fat in, 402 manufacture of, 393 methods of, analysis, 398 microscopical structure, 403 nibs, 394, 402 nitrogeneous bodies in, 396 pentosans in, 396 references on, 406 shells, 394, 395, 405 standards, 402 sugar in, 390, 405 sugar in, 390, 405 sugar in, 390, 405 sugar in, 390, 405 sugar in, 390, 405 choromine in, 396, 400 Cocoanut oil, 528 pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 384 caffeine in, 380, 384 caffeine in, 370 chicory in, 388, 389 coloring of, 384 composition of, 379, 380, 381 essential oil of, 379 fat in, 379 glazing of, 385 microscopical structure, 386 "pellets," 384 microscopical structure, 386 "pellets," 384 microscopical structure, 386 "pellets," 384	dyeing wool by, 793	Cognac, 739. See also Brandy.
identification of, 793, 797, 799, 803 in milk, 177 in sausages, 239 Rota's scheme for, 797 Sostegni and Carpentieri's color test, 794 separation with ether, 796 Cochineal, 790 in sausages, 238 Cocoa, 392, 393 adulteration of, 402 alkali in, 403 ash of, 396 butter, 393, 590 caffeine in, 402 manufacture of, 393 methods of, analysis, 398 microscopical structure, 403 nibs, 394, 402 nitrogeneous bodies in, 396 pentosans in, 396 references on, 406 shells, 394, 395, 405 standards, 402 starch in, 394, 395, 495 standards, 402 sugar in, 399, 405 theobromine in, 396, 400 Cocoanut oil, 528 pulp, 528 shells, 410 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffeine in, 380, 384 caffetannic acid in, 379, 382 caffeine in, 379 chicory in, 388, 389 coloring of, 384 composition of, 379, 380, 381 essential oil of, 379 fat in, 579 glazing of, 385 microscopical structure, 386 "pellets," 384 microscopical structure, 386 "pellets," 374 milk, 177 Colorometric analysis, 77 Colorometric analysis, 77 Colorometric analysis, 77 Colorometric analysis, 77 Colorometric analysis, 77 Colorometric analysis, 77 Colorometric analysis, 77 Colorometric analysis, 77 Colorometric analysis, 77 Colorometric analysis, 78 alci fuchsin, 790 arisenic compounds, 782 barium compounds, 782 basic, 793 blue, 783, 784, 786, 792, 810 brown, 784, 786, 808 caramel, 790 coal tar, 791, 792, 793 cochineal, 790 coal tar, 791, 792, 793 cochineal, 790 coal tar, 791, 792, 793 cochineal, 790 coal tar, 791, 792, 793 cochineal, 790 coal tar, 791, 792, 793 cochineal, 790 coal tar, 791, 792, 793 cochineal, 790 coal tar, 791, 792, 793 cochineal, 790 coal tar, 791, 792, 793 cochineal, 790 coal tar, 791, 792, 793 cochineal, 790 coal tar, 791, 792, 793 cochineal, 790 coal tar, 791, 792, 793 cochineal, 790 coal tar, 791, 792, 793 cochineal, 790 coal tar, 791, 792, 793 cochineal, 790 coal tar, 791, 792, 793 cochineal, 790 coal tar, 791, 792, 793 cochineal, 790 coal tar, 791, 792, 793 cochineal, 790 coal tar, 791, 793 spandium compounds, 782 gramely, 790 coal tar, 791, 792, 793 c	extraction by amyl alcohol,	oil, 741
799, 803 in milk, 177 in sausages, 239 Rota's scheme for, 797 Sostegan and Carpentieri's color test, 794 separation with ether, 796 Cochineal, 790 in sausages, 238 Cocoa, 392, 393 adulteration of, 402 alkali in, 403 ash of, 396 butter, 393, 529 caffeine in, 400 composition of, 393 foreign fat in, 402 manufacture of, 393 methods of, analysis, 398 microscopical structure, 403 nibs, 394, 402 nitrogeneous bodies in, 396 pentosans in, 396 pentos	795	Collagen, 42, 211
in milk, 177 in sausages, 239 Rota's scheme for, 797 Sostegni and Carpentieri's color test, 794 separation with ether, 796 Cochineal, 790 in sausages, 238 Cocoa, 392, 393 adulteration of, 402 alkali in, 403 ash of, 396 butter, 393, 529 caffeine in, 402 manufacture of, 393 methods of, analysis, 398 microscopical structure, 403 nibs, 394, 402 nitrogeneous bodies in, 396 pentosans in, 396 references on, 406 shells, 394, 395, 405 standards, 402 starch in, 394, 495 sugar in, 399, 405 theobromine in, 396, 400 Cocoanut oil, 528 pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffetin in, 390, 393 cochineal, 790 copper compounds, 782 coal tar, 791, 792, 793 cochineal, 790 copper compounds, 782 coal tar, 791, 792, 793 cochineal, 790 copper compounds, 782 coal tar, 791, 792, 793 cochineal, 790 copper compounds, 782 coal tar, 791, 792, 793 cochineal, 790 copper compounds, 782 coloring, 789, 784, 786, 808 caramel, 790 cochineal, 790 composition of, 393 methods of, 393 methods of, 384 composition of, 379 fat in, 379 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384 Colormetric analysis, 77 Colors, artificial, 780 acid fuchsin 799 allowed, 792 animal, 790 arsenic compounds, 783 barium compounds, 782 basic, 793 blue, 783, 784, 786, 808 caramel, 790 cool tar, 792, 793 cochineal, 790 copper compounds, 782 coal tar, 791, 792, 793 cochineal, 790 copper compounds, 782 coal tar, 791, 792, 793 cochineal, 790 copper compounds, 782 coal tar, 791, 792, 793 cochineal, 790 copper compounds, 782 coal tar, 796, 808 caramel, 790 copper compounds, 782 coloring, 784, 786, 808 caramel, 790 colotineal, 790 copper compounds, 782 cothera, 786, 808 caramel, 790 colotineal, 790 copper compounds, 782 cothera, 786, 808 caramel, 790 colotineal, 790 copper compounds, 782 cothera, 786, 808 caramel, 790 colotineal, 790 copper compounds, 782 cothera, 786, 808 caramel, 790 colotineal, 790 copper compounds, 782 cothera, 786, 808 caramel, 790 coherate, 785, 784, 786, 789, 810 in mile,	identification of, 793, 797,	Collodion silk, 705
in sausages, 239 Rota's scheme for, 797 Sostegni and Carpentieri's color test, 794 separation with ether, 796 Cochineal, 790 in sausages, 238 Cocoa, 392, 393 adulteration of, 402 alkali in, 403 ash of, 396 butter, 393, 529 caffeine in, 400 composition of, 393 foreign fat in, 402 manufacture of, 393 methods of, analysis, 398 microscopical structure, 403 nibs, 394, 402 nitrogeneous bodies in, 396 pentosans in, 396 references on, 406 shells, 394, 395, 495 standards, 402 starch in, 394, 395, 399, 405 sugar in, 399, 405 sugar in, 399, 405 theobromine in, 396, 400 Cocoanut oil, 528 pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffeine in, 380, 384 caffetannic acid in, 379, 382 caffeol in, 379 chicory in, 388, 389 coloring of, 384 composition of, 379, 380, 381 essential oil of, 379 fat in, 379 glazing of, 384 composition of, 379, 380, 381 essential oil of, 379 fat in, 579 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384	799, 803	Colorimeter, Schreiner's, 77
Rota's scheme for, 797 Sostegni and Carpentieri's color test, 794 separation with ether, 796 Cochineal, 790 in sausages, 238 Cocoa, 392, 393 adulteration of, 402 alkali in, 403 ash of, 396 butter, 393, 529 caffeine in, 400 composition of, 393 foreign fat in, 402 manufacture of, 393 methods of, analysis, 398 microscopical structure, 403 nibs, 394, 402 nitrogeneous bodies in, 396 pentosans in, 396 references on, 406 shells, 394, 395, 405 standards, 402 starch in, 394, 395, 399, 405 sugar in, 399, 405 theobromine in, 396, 400 Cocoanut oil, 528 pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffeine in, 380, 384 caffetannic acid in, 379, 380 ash of, 380, 382 caffeine in, 380, 384 caffetannic acid in, 379, 380, 381 essential oil off, 379 fat in, 579 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384	in milk, 177	Colorometric analysis, 77
Sostegni and Carpentieri's color test, 794 separation with ether, 796 Cochineal, 790 in sausages, 238 Cocoa, 392, 393 adulteration of, 402 alkali in, 403 ash of, 396 butter, 393, 529 caffeine in, 400 composition of, 393 foreign fat in, 402 manufacture of, 393 methods of, analysis, 398 microscopical structure, 403 nibs, 394, 402 nitrogeneous bodies in, 396 pentosans in, 396 references on, 406 shells, 394, 395, 405 standards, 402 starch in, 3904, 495 theobromine in, 396, 400 Cocoanut oil, 528 pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffetannic acid in, 379, 382 caffetannic acid in, 379, 382 caffetannic acid in, 379, 382 caffetannic acid in, 379, 380, 381 essential oil of, 379 fat in, 579 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384 and lateration compounds, 783 barium compounds, 782 basic, 793 blue, 783, 784, 786, 792, 810 brown, 784, 786, 808 caramel, 790 col tar, 790, 792, 793 cochinal, 790 col tar, 790, 792, 810 brown, 784, 786, 808 caramel, 790 col tar, 790, 792, 810 brown, 784, 786, 808 caramel, 790 col tar, 790, 792, 810 brown, 784, 786, 808 caramel, 790 colatar, 790 colatar, 790 colatar, 795 fuchsin, 796 greer, 783, 784, 786, 792, 810 brown, 784, 786, 808 caramel, 790 colatar, 790 colatar, 790 colatar, 790 colatar, 790 colatar, 795 fuchsin, 798 green, 783, 784, 786, 792, 810 brown, 784, 786, 808 caramel, 790 colatar, 790 colatar, 790 colatar, 790 colatar, 795 fuchsin, 798 green, 783, 784, 786, 792, 810 brown, 784, 786, 808 caramel, 790 colatar, 790 colatar, 790 colatar, 790 colatar, 790 colatar, 795 fuchsin, 798 green, 783, 784, 786, 792, 810 brown, 784, 786, 808 caramel, 790 colatar, 790 colatar, 790 colatar, 790 colatar, 790 colatar, 790 colatar, 790 colatar, 790 colatar, 790 colatar, 790 colatar, 790 colatar, 790 colatar, 790 colatar, 780, 792, 810 harmless, 782, 784 in butter, 335 in butter, 335 in butter, 337 in		
color test, 794 separation with ether, 796 Cochineal, 790 in sausages, 238 Cocoa, 392, 393 adulteration of, 402 alkali in, 403 ash of, 396 butter, 393, 529 caffeine in, 400 composition of, 393 methods of, analysis, 382 microscopical structure, 403 nibs, 394, 402 nitrogeneous bodies in, 396 pentosans in, 396 references on, 406 shells, 394, 395, 495 standards, 402 starch in, 394, 495 sugar in, 399, 495 theobromine in, 396, 400 Cocoanut oil, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffeine in, 380, 384 caffetannic acid in, 379, 382 caffetol in, 379 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384 animal, 790 arsenic compounds, 782 basic, 793 blue, 783, 784, 786, 792, 810 brown, 784, 786, 808 caramel, 790 cool tar, 791, 792, 793 cochineal, 790 copper compounds, 782 codhieal, 790 copper compounds, 782 cochineal, 790 copper compounds, 782 cochineal, 790 copper compounds, 782 cochineal, 790 copper compounds, 782 cudbear, 789 extraction of, by immiscible solvents, 795 fuchsin, 798 green, 783, 784, 786, 792, 810 brown, 784, 786, 808 caramel, 790 copper compounds, 782 cudbear, 789 extraction of, by immiscible solvents, 795 fuchsin, 798 green, 783, 784, 785, 792, 810 harmless, 782, 783, 784, 785, 792, 810 harmless, 782, 783, 784, 785, 792, 810 harmless, 782, 783, 784, 785, 792, 810 harmless, 782, 784, 785, 792, 810 harmless, 782, 783, 784, 785, 792, 810 harmless, 782, 783, 784, 785, 792, 810 harmless, 782 fuchsin, 798 green, 783, 784, 785, 792, 810 harmless, 782 fuchsin, 798 inductor, 783 fuchsia, 796 composition of, 393 metrodos of, 394 incomposition of, 393 inductore of		
Cochineal, 790		
Cochineal, 790 in sausages, 238 Cocoa, 392, 393 adulteration of, 402 alkali in, 403 ash of, 396 butter, 393, 529 caffeine in, 400 composition of, 393 methods of, analysis, 398 microscopical structure, 403 nibs, 394, 402 nitrogeneous bodies in, 396 pentosans in, 396 references on, 406 shells, 394, 395, 405 starch in, 394, 395, 399, 405 sugar in, 399, 405 theobromine in, 396, 400 Cocoanut oil, 528 pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffeine in, 390, 384 caffetannic acid in, 379, 582 caffein in, 379 chicory in, 388, 389 coloring of, 384 composition of, 379, 380, 381 essential oil of, 379 fat in, 579 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 403 miss and jellies, 921 in ketchup, 908 in milk, 174-177 in mustard, 460 in sugar, 590 lead chromate, 647, 782 lead compounds, 782 basic, 793 brown, 784, 786, 808 caramel, 790 cohromate of lead, 791 coal tar, 791, 792, 793 cochineal, 790 cohromate of lead, 791 coal tar, 791, 792, 793 cochineal, 790 copper compounds, 782 cudbear, 789 extraction of, 59 fuchsin, 798 green, 783, 784, 786, 808 caramel, 790 copper compounds, 782 cudbear, 789 extraction of, 59 fuchsin, 795 fuchsin, 795 fuchsin, 795 fuchsin, 796 fuchsin, 796 span, 783, 784, 786, 808 caramel, 790 cohromate of lead, 791 coal tar, 791, 792, 793 cochineal, 790 cohromate of lead, 791 coal tar, 791, 792, 793 cochineal, 790 cohromate of lead, 791 coal tar, 791, 792, 793 cochineal, 790 cohromate of lead, 791 coal tar, 791, 792, 793 cochineal, 790 cohromant, 782 interation of, 93 petraction of, 94 petraction of, 94 pe		
Cocoa, 392, 393 adulteration of, 402 alkali in, 403 ash of, 396 butter, 393, 529 caffeine in, 400 composition of, 393 foreign fat in, 402 manufacture of, 393 methods of, analysis, 398 microscopical structure, 403 nibs, 394, 402 nitrogeneous bodies in, 396 pentosans in, 396 references on, 406 shells, 394, 395, 405 standards, 402 starch in, 394, 395, 405 standards, 402 starch in, 394, 495 theobromine in, 396, 400 Cocoanut oil, 528 pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffeine in, 380, 384 caffetannic acid in, 379, 582 caffeol in, 379 chicory in, 388, 389 coloring of, 384 composition of, 379, 380, 381 essential oil of, 379 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384		
Cocoa, 392, 393 adulteration of, 402 alkali in, 403 ash of, 396 butter, 393, 529 caffeine in, 400 composition of, 393 foreign fat in, 402 manufacture of, 393 methods of, analysis, 398 microscopical structure, 403 nibs, 394, 402 nitrogeneous bodies in, 396 pentosans in, 396 references on, 406 shells, 394, 395, 405 standards, 402 starch in, 394, 395, 405 standards, 402 starch in, 394, 395, 399, 405 sugar in, 399, 405 theobromine in, 396, 400 Cocoanut oil, 528 pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffeine in, 380, 384 caffetannic acid in, 379, 582 caffeol in, 379 chicory in, 388, 389 coloring of, 384 composition of, 379 fat in, 379 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384	1.77	
adulteration of, 402 alkali in, 403 ash of, 396 butter, 393, 529 caffeine in, 400 composition of, 393 foreign fat in, 402 manufacture of, 393 methods of, analysis, 398 microscopical structure, 403 nibs, 394, 402 nitrogeneous bodies in, 396 pentosans in, 396 references on, 406 shells, 394, 395, 405 standards, 402 starch in, 394, 395, 399, 405 sugar in, 399, 405 sugar in, 399, 405 sugar in, 399, 405 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffein in, 380, 384 caffetannic acid in, 379, 582 caffeol in, 379 chicory in, 388, 389 coloring of, 384 composition of, 379, 380, 381 essential oil of, 379 fat in, 379 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384 brown, 784, 786, 808 caramel, 790 cochicnat, 790 cochical, 790 copper compounds, 782 cudbear, 789 extraction of, by immiscible solvents, 795 fuchsin, 798 green, 783, 784, 785, 792, 810 harmless, 782, 784, 785, 792, 810 harmless, 782, 784, 785, 792, 810 harmless, 782, 784, 785, 792, 810 injurious, 782, 783 in butter, 336 in digo, 790, 810 injurious, 782, 783 in butter, 336 in milk, 174–177 in mustard, 460 in sugar, 590 lead chromate, 647, 782 lead compounds, 782 logenter, 783, 784, 785, 792, 810 injurious, 782, 783 in butter, 336 in butter, 335 in cayenne, 444 in confectionery, 649, 786 in jams and jellies, 921 in ketchup, 908 in milk, 174–177 in mustard, 460 in sugar, 590 lead chromate, 647, 782 lead compounds, 782 logenter, 783, 784, 785, 792, 810 injurious, 782, 783 in butter, 336 in butter, 336 in butter, 336 in injurious, 782, 783 in butter, 335 in cayenne, 444 in confectionery, 649, 786 in jams and jellies, 921 in ketchup, 908 in milk, 174–177 in mustard, 460 in sugar, 590 lead chromate, 647, 782 lead compounds, 782 logenter, 783, 784, 785, 792, 810 injurious, 782, 783 in butter, 335 in cayenne, 444 in confectionery, 649, 786 in jams and jellies, 921 in ketchup, 908 in milk, 174–177 in mustard, 460 in sugar, 590 lead chromate, 647, 782 lead compounds, 782 logenter, 783, 784, 785, 792, 81		
alkali in, 403 ash of, 396 butter, 393, 529 caffeine in, 400 composition of, 393 foreign fat in, 402 manufacture of, 393 methods of, analysis, 398 microscopical structure, 403 nibs, 394, 402 nitrogeneous bodies in, 396 pentosans in, 396 references on, 406 shells, 394, 395, 405 standards, 402 starch in, 399, 405 sugar in, 399, 405 sugar in, 399, 405 sugar in, 399, 405 consenut oil, 528 pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffein in, 360, 384 caffetannic acid in, 379, 582 caffeol in, 379 chicory in, 388, 389 coloring of, 384 composition of, 379, 380, 381 essential oil of, 379 flat in, 579 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384 caffet of, 781 coal tar, 791, 792, 793 cochineal, 790 copper compounds, 782 cudbear, 795 fuchsin, 796 green, 783, 784, 785, 792, 810 harmless, 782, 784 identification of, 793, 797, 803 indigo, 790, 810 injurious, 782, 783 in butter, 535 in cayenne, 444 in confectionery, 649, 786 in jams and jellies, 921 in ketchup, 908 in milk, 174–177 in mustard, 460 in sugar, 590 lead chromate, 647, 782 lead compounds, 782 loghes, 789, 780 mercury compounds, 782 ochineal, 790 copper compounds, 782 cudbear, 791, 792, 810 harmless, 782, 784 identification of, 793, 797, 803 indigo, 790, 810 in jurious, 782, 783 in butter, 535 in cayenne, 444 in confectionery, 649, 786 in jams and jellies, 921 in ketchup, 908 in milk, 174–177 in mustard, 460 in sugar, 590 lead chromate, 647, 782 lead compounds, 782 cudbear, 791, 803 indigo, 790, 810 injurious, 782, 783 in butter, 535 in cayenne, 444 in confectionery, 649, 786 in jams and jellies, 921 in ketchup, 908 in milk, 174–177 in mustard, 460 in sugar, 590 lead chromate, 647, 782 lead compounds, 782 lead compounds, 782 lead compounds, 782 lead compounds, 782 loghestic, 798 mercury compounds, 782 loghestic, 798 indigo, 790, 810 inquious, 782, 784 in confectionery, 649, 786 in jams and jellies, 921 in ketchup, 908 in milk, 174–177 in mustard, 460 in sugar, 590 lead chromate,		
ash of, 396 butter, 393, 529 caffeine in, 400 composition of, 393 foreign fat in, 402 manufacture of, 393 methods of, analysis, 398 microscopical structure, 403 nibs, 394, 402 nitrogeneous bodies in, 396 pentosans in, 396 references on, 406 shells, 394, 395, 405 standards, 402 starch in, 394, 395, 399, 405 theobromine in, 396, 400 Cocoanut oil, 528 pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffein in, 380, 384 caffetannic acid in, 379, 382 caffeol in, 379 chicory in, 388, 389 coloring of, 384 composition of, 379, 380, 381 essential oil of, 379 fat in, 579 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384		
butter, 393, 529 caffeine in, 400 composition of, 393 foreign fat in, 402 manufacture of, 393 methods of, analysis, 398 microscopical structure, 403 nibs, 394, 402 nitrogeneous bodies in, 396 pentosans in, 396 references on, 406 shells, 394, 395, 405 standards, 402 starch in, 390, 405 sugar in, 399, 405 stells, 419 Cocoanut oil, 528 pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffeine in, 390, 384 composition of, 379, 380, 381 essential oil of, 379 fat in, 579 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384		* * * *
caffeine in, 400 composition of, 393 foreign fat in, 402 manufacture of, 393 methods of, analysis, 398 microscopical structure, 403 nibs, 394, 402 nitrogeneous bodies in, 396 pentosans in, 396 references on, 406 shells, 394, 395, 405 standards, 402 starch in, 394, 395, 399, 405 sugar in, 399, 405 theobromine in, 396, 400 Cocoanut oil, 528 pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffein in, 380, 384 caffetannic acid in, 379, 582 caffein in, 388, 389 coloring of, 384, 389 coloring of, 384, 389 coloring of, 384, 389 coloring of, 384, 389 coloring of, 384, 389 coloring of, 384, 389 coloring of, 384, 389 coloring of, 384, 389 coloring of, 384, 389 coloring of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384		
composition of, 393 foreign fat in, 402 manufacture of, 393 methods of, analysis, 398 microscopical structure, 403 nibs, 394, 402 nitrogeneous bodies in, 396 pentosans in, 396 references on, 406 shells, 394, 395, 405 standards, 402 starch in, 394, 395, 399, 405 sugar in, 399, 405 theobromine in, 396, 400 Cocoanut oil, 528 pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffeine in, 380, 384 caffetannic acid in, 379, 382 caffetol in, 379 chicory in, 388, 389 coloring of, 384 composition of, 379, 380, 381 essential oil of, 379 fat in, 579 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384		
foreign fat in, 402 manufacture of, 393 methods of, analysis, 398 microscopical structure, 403 nibs, 394, 402 nitrogeneous bodies in, 396 pentosans in, 396 references on, 406 shells, 394, 395, 405 standards, 402 starch in, 394, 395, 399, 405 sugar in, 399, 405 theobromine in, 396, 400 Cocoanut oil, 528 pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffein in, 380, 384 caffetannic acid in, 379, 582 caffetol in, 379 chicory in, 388, 389 coloring of, 384 composition of, 379, 380, 381 essential oil of, 379 fat in, 579 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384 cudbear, 789 extraction of, by immiscible solvents, 795 fuchsin, 798 green, 783, 784, 785, 792, 810 harmless, 782, 784 identification of, 793, 797, 803 indigo, 790, 810 injurious, 782, 783 in butter, 535 in cayenne, 444 in confectionery, 649, 786 in jams and jellies, 921 in ketchup, 908 in milk, 174–177 in mustard, 460 in sugar, 590 lead chromate, 647, 782 lead compounds, 782 lead compounds, 782 lead compounds, 782 lead compounds, 782 lead compounds, 782 lead compounds, 782 orange, 785, 792, 806 orchil, 789 Prussian blue, 790, 792, 810 reagents for identifying, 812 red, 783, 784, 788 references on, 813 Rota's scheme for, 797 separation by solvents, 795 toxic effect of, 781		
manufacture of, 393 methods of, analysis, 398 microscopical structure, 403 nibs, 394, 402 nitrogeneous bodies in, 396 pentosans in, 396 references on, 406 shells, 394, 395, 405 standards, 402 starch in, 394, 395, 399, 405 stagar in, 399, 405 theobromine in, 396, 400 Cocoanut oil, 528 pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffein in, 380, 384 caffetannic acid in, 379, 382 caffeol in, 379 chicory in, 388, 389 coloring of, 384 composition of, 379 fat in, 579 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384 extraction of, by immiscible solvents, 795 fuchsin, 798 green, 783, 784, 785, 792, 810 harmless, 782, 784 identification of, 793, 797, 803 indigo, 790, 810 injurious, 782, 783 in butter, 535 in cayenne, 444 in confectionery, 649, 786 in jams and jellies, 921 in ketchup, 908 in milk, 174-177 in mustard, 460 in sugar, 590 lead chromate, 647, 782 lead compounds, 782 lead compounds, 782 logwood, 789 mercury compounds, 783 mineral, 790 non-injurious, 782, 784 orange, 785, 792, 806 orchil, 789 Prussian blue, 790, 792, 810 reagents for identifying, 812 red, 783, 784, 788, 792, 810 harmless, 782, 784 identification of, 793, 797, 803 indigo, 790, 810 in jurious, 782, 784 in confectionery, 649, 786 in jams and jellies, 921 in ketchup, 908 in milk, 174-177 in mustard, 460 in sugar, 590 lead chromate, 647, 782 lead compounds, 782 lead compounds, 782 lead compounds, 782 lead compounds, 782 lead compounds, 782 lead compounds, 782 lead compounds, 782 lead compounds, 782 lead compounds, 782 lead compounds, 782 lead compounds, 782 lead compounds, 782 lead compounds, 782 lead compounds, 782 lead compounds, 782 lead compounds, 782 lead compounds, 783 mineral, 790 non-injurious, 782, 784 orangene, 444 in confectionery, 649, 786 in jurious, 782, 784 orangene, 444 in confectionery, 649, 786 in jurious, 782, 784 orangene, 444 in confectionery in mustard, 460 in sugar, 590 lead chromate, 647, 782 lead compounds, 782 lead compounds, 783 mineral, 790 n		• • • • • • • • • • • • • • • • • • • •
methods of, analysis, 398 microscopical structure, 403 nibs, 394, 402 nitrogeneous bodies in, 396 pentosans in, 396 references on, 406 shells, 394, 395, 495 standards, 402 starch in, 394, 395, 399, 405 stugar in, 399, 405 theobromine in, 396, 400 Cocoanut oil, 528 pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffein in, 380, 384 caffetannic acid in, 379, 382 caffeol in, 379 chicory in, 388, 389 coloring of, 384 composition of, 379, 380, 381 essential oil of, 379 fat in, 379 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 403 fuchsin, 798 green, 783, 784, 785, 792, 810 harmless, 782, 784 identification of, 793, 797, 803 indigo, 790, 810 injurious, 782, 783 in butter, 535 in cayenne, 444 in confectionery, 649, 786 in jams and jellies, 921 in ketchup, 908 in milk, 174-177 in mustard, 460 in sugar, 590 lead chromate, 647, 782 lead compounds, 782 logwood, 789 mercury compounds, 783 mineral, 790 non-injurious, 782, 784 orange, 785, 792, 810 reagents for identifying, 812 red, 783, 784, 788 references on, 813 Rota's scheme for, 797 separation by solvents, 795 toxic effect of, 781	~	
microscopical structure, 403 nibs, 394, 402 nitrogeneous bodies in, 396 pentosans in, 396 references on, 406 shells, 394, 395, 405 standards, 402 starch in, 394, 395, 399, 405 sugar in, 399, 405 theobromine in, 396, 400 Cocoanut oil, 528 pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffein in, 380, 384 caffetannic acid in, 379, 582 caffeol in, 379 chicory in, 388, 389 coloring of, 384 composition of, 379, 380, 381 essential oil of, 379 fat in, 379 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384 "pellets," 384 fuchsin, 798 green, 783, 784, 785, 792, 810 harmless, 782, 784 identification of, 793, 797, 803 indigo, 790, 810 injurious, 782, 783 in butter, 535 in cayenne, 444 in confectionery, 649, 786 in jams and jellies, 921 in ketchup, 908 in milk, 174-177 in mustard, 460 in sugar, 590 lead chromate, 647, 782 lead compounds, 782 lead compounds, 783 mineral, 790 non-injurious, 782, 784 orange, 785, 792, 806 orchil, 789 Prussian blue, 790, 792, 810 reagents for identifying, 812 red, 783, 784, 788, 784, 788 references on, 813 Rota's scheme for, 797 separation by solvents, 795 toxic effect of, 781		· · · · · · · · · · · · · · · · · · ·
nibs, 394, 402 nitrogeneous bodies in, 396 pentosans in, 396 pentosans in, 396 references on, 406 shells, 394, 395, 405 standards, 402 starch in, 394, 395, 399, 405 sugar in, 399, 405 theobromine in, 396, 400 Cocoanut oil, 528 pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffeine in, 380, 384 caffetannic acid in, 379, 582 caffeol in, 379 chicory in, 388, 389 coloring of, 384 composition of, 379, 380, 381 essential oil of, 379 fat in, 579 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384 "pellets," 384 "pellets," 384 "pellets," 384 "pellets," 384 "pellets," 384 "green, 783, 784, 785, 792, 810 harmless, 782, 784 identification of, 793, 797, 803 indigo, 790, 810 injurious, 782, 783 in butter, 535 in cayenne, 444 in confectionery, 649, 786 in jams and jellies, 921 in ketchup, 908 in milk, 174–177 in mustard, 460 in sugar, 590 lead chromate, 647, 782 lead compounds, 782 lead compounds, 783 mineral, 790 non-injurious, 782, 784 orange, 785, 792, 806 orchil, 789 Prussian blue, 790, 792, 810 reagents for identifying, 812 red, 783, 784, 785, 792, 803 indigo, 790, 810 indigo, 790, 810 indigo, 790, 810 indigo, 790, 810 indigo, 790, 810 indigo, 790, 810 indigo, 790, 810 indigo, 790, 810 in jurious, 782, 783 in butter, 535 in cayenne, 444 in confectionery, 649, 786 in jams and jellies, 921 in ketchup, 908 in milk, 174–177 in mustard, 460 in sugar, 590 lead chromate, 647, 782 lead compounds, 783 mineral, 790 non-injurious, 782, 784 orangent, 783 in butter, 535 in cayenne, 444 in confectionery, 649, 786 in jams and jellies, 921 in ketchup, 908 in milk, 174–177 in mustard, 460 in sugar, 590 lead chromate, 647, 782 lead compounds, 782 lead compounds, 783 mineral, 790 non-injurious, 782, 784 orangent, 783 in butter, 535 in cayenne, 444 in confectionery in cayenne, 444 in confectionery in cayenne, 444 in confectionery in users, 790 in mustard, 460 in sugar, 590 lead chromate, 647, 782 lead compounds, 783 mineral, 790 reader defense of 782 in mustar		
nitrogeneous bodies in, 396 pentosans in, 396 references on, 406 shells, 394, 395, 405 standards, 402 starch in, 394, 395, 399, 405 sugar in, 399, 405 theobromine in, 396, 400 Cocoanut oil, 528 pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffeine in, 380, 384 caffetannic acid in, 379, 382 caffeol in, 379 chicory in, 388, 389 coloring of, 384 composition of, 379, 380, 381 essential oil of, 379 fat in, 579 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384 harmless, 782, 784 identification of, 793, 797, 803 indigo, 790, 810 injurious, 782, 783 in butter, 535 in cayenne, 444 in confectionery, 649, 786 in jams and jellies, 921 in ketchup, 908 in milk, 174-177 in mustard, 460 in sugar, 590 lead chromate, 647, 782 lead compounds, 782 lead compounds, 782 microscopical structure, 386 harmless, 782, 784 identification of, 793, 797, 803 indigo, 790, 810 injurious, 782, 783 in butter, 535 in cayenne, 444 in confectionery, 649, 786 in jams and jellies, 921 in ketchup, 908 in milk, 174-177 in mustard, 460 in sugar, 590 lead chromate, 647, 782 lead compounds, 782 lead compounds, 783 mineral, 790 ornoi-injurious, 782, 784 orange, 785, 792, 806 orchil, 789 Prussian blue, 790, 792, 810 reagents for identifying, 812 red, 783, 784, 788 references on, 813 Rota's scheme for, 797 separation by solvents, 795 toxic effect of, 781	- 1	
pentosans in, 396 references on, 406 shells, 394, 395, 405 standards, 402 starch in, 394, 395, 399, 405 sugar in, 399, 405 theobromine in, 396, 400 Cocoanut oil, 528 pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffeine in, 380, 384 caffetannic acid in, 379, 382 caffeol in, 379 chicory in, 388, 389 coloring of, 384 composition of, 379, 380, 381 essential oil of, 379 fat in, 579 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 identification of, 793, 797, 803 indigo, 790, 810 injurious, 782, 783 in butter, 535 in cayenne, 444 in confectionery, 649, 786 in jams and jellies, 921 in ketchup, 908 in milk, 174–177 in mustard, 460 in sugar, 590 lead chromate, 647, 782 lead compounds, 782 logwood, 789 mercury compounds, 783 mineral, 790 ono-injurious, 782, 784 orange, 785, 792, 806 orchil, 789 Prussian blue, 790, 792, 810 reagents for identifying, 812 red, 783, 784, 788 references on, 813 Rota's scheme for, 797 separation by solvents, 795 toxic effect of, 781		
references on, 406 shells, 394, 395, 405 standards, 402 starch in, 394, 395, 399, 405 sugar in, 399, 405 theobromine in, 396, 400 Cocoanut oil, 528 pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffeine in, 380, 384 caffetannic acid in, 379, 582 caffeol in, 379 chicory in, 388, 389 coloring of, 384 composition of, 379, 380, 381 essential oil of, 379 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384 in butter, 535 in cayenne, 444 in confectionery, 649, 786 in jams and jellies, 921 in ketchup, 908 in milk, 174–177 in mustard, 460 in sugar, 590 lead chromate, 647, 782 lead compounds, 782 logwood, 789 mercury compounds, 783 mineral, 790 non-injurious, 782, 784 orange, 785, 792, 806 orchil, 789 Prussian blue, 790, 792, 810 reagents for identifying, 812 red, 783, 784, 788 references on, 813 Rota's scheme for, 797 separation by solvents, 795 toxic effect of, 781		
shells, 394, 395, 405 standards, 402 starch in, 394, 395, 399, 405 sugar in, 399, 405 theobromine in, 396, 400 Cocoanut oil, 528 pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffeine in, 379 chicory in, 388, 389 coloring of, 384 coloring of, 384 composition of, 379, 380, 381 essential oil of, 379 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384 in jurious, 782, 783 in butter, 535 in cayenne, 444 in confectionery, 649, 786 in jams and jellies, 921 in ketchup, 908 in milk, 174-177 in mustard, 460 in sugar, 590 lead chromate, 647, 782 lead compounds, 782 logwood, 789 mercury compounds, 783 mineral, 790 non-injurious, 782, 784 orange, 785, 792, 806 orchil, 789 Prussian blue, 790, 792, 810 reagents for identifying, 812 references on, 813 Rota's scheme for, 797 separation by solvents, 795 toxic effect of, 781		
standards, 402 starch in, 394, 395, 399, 405 sugar in, 399, 405 theobromine in, 396, 400 Cocoanut oil, 528 pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffeine in, 380, 384 caffetannic acid in, 379, 582 caffeol in, 379 chicory in, 388, 389 coloring of, 384 composition of, 379, 380, 381 essential oil of, 379 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384		
starch in, 394, 395, 399, 405 sugar in, 399, 405 theobromine in, 396, 400 Cocoanut oil, 528 pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffeine in, 380, 384 caffetannic acid in, 379, 382 caffeol in, 379 chicory in, 388, 389 coloring of, 384 composition of, 379, 380, 381 essential oil of, 379 fat in, 579 glazing of, 385 hygienic, 390 methods of analysis, 382 methods of analysis, 382 methods of analysis, 382 methods of analysis, 382 methods of analysis, 386 "pellets," 384 in cayenne, 444 in confectionery, 649, 786 in jams and jellies, 921 in ketchup, 908 in milk, 174-177 in mustard, 460 in sugar, 590 lead chromate, 647, 782 lead compounds, 782 lead compounds, 783 mercury compounds, 783 mineral, 790 non-injurious, 782, 784 orange, 785, 792, 806 orchil, 789 Prussian blue, 790, 792, 810 reagents for identifying, 812 red, 783, 784, 788 references on, 813 Rota's scheme for, 797 separation by solvents, 795 toxic effect of, 781		
sugar in, 399, 405 theobromine in, 396, 400 Cocoanut oil, 528 pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffeine in, 380, 384 caffetannic acid in, 379, 382 caffeol in, 379 chicory in, 388, 389 coloring of, 384 composition of, 379, 380, 381 essential oil of, 379 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 in confectionery, 649, 786 in jams and jellies, 921 in ketchup, 908 in milk, 174-177 in mustard, 460 in sugar, 590 lead chromate, 647, 782 lead compounds, 782 lead compounds, 782 mercury compounds, 783 mineral, 790 non-injurious, 782, 784 orange, 785, 792, 806 orchil, 789 Prussian blue, 790, 792, 810 reagents for identifying, 812 red, 783, 784, 788 references on, 813 Rota's scheme for, 797 separation by solvents, 795 toxic effect of, 781		- · · · · · · · · · · · · · · · · · · ·
theobromine in, 396, 400 Cocoanut oil, 528 pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffeine in, 380, 384 caffetannic acid in, 379, 382 caffeol in, 379 chicory in, 388, 389 coloring of, 384 composition of, 379, 380, 381 essential oil of, 379 fat in, 379 glazing of, 385 hygienic, 390 methods of analysis, 382 methods of analysis, 382 methods of analysis, 382 methods of analysis, 382 methods of analysis, 386 "pellets," 384 in ketchup, 908 in milk, 174–177 in mustard, 460 in sugar, 590 lead chromate, 647, 782 lead compounds, 782 lead compounds, 782 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 782 lead compounds, 782 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 782 lead compou		
Cocoanut oil, 528 pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffein in, 380, 384 caffetannic acid in, 379, 382 caffeol in, 379 chicory in, 388, 389 coloring of, 384 composition of, 379, 380, 381 essential oil of, 379 fat in, 379 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384 in ketchup, 908 in milk, 174–177 in mustard, 460 in sugar, 590 lead chromate, 647, 782 lead compounds, 782 lead compounds, 782 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 783 mercury compounds, 782 references on, 813 Rota's scheme for identifying, 812 red, 783, 784, 788 references on, 813 Rota's scheme for, 797 separation by solvents, 795 toxic effect of, 781		
pulp, 528 shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffeine in, 380, 384 caffetannic acid in, 379, 382 caffeol in, 379 chicory in, 388, 389 coloring of, 384 composition of, 379, 380, 381 essential oil of, 379 fat in, 379 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384 in milk, 174–177 in mustard, 460 in sugar, 590 lead chromate, 647, 782 lead compounds, 782 lead compounds, 782 mercury compounds, 783 mineral, 790 range, 785, 792, 806 orchil, 789 Prussian blue, 790, 792, 810 reagents for identifying, 812 red, 783, 784, 788 references on, 813 Rota's scheme for, 797 separation by solvents, 795 toxic effect of, 781		
shells, 419 Coffee, 379 adulteration of, 384 ash of, 380, 382 caffeine in, 380, 384 caffetannic acid in, 379, 582 caffeol in, 379 chicory in, 388, 389 coloring of, 384 composition of, 379, 380, 381 essential oil of, 379 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384 in mustard, 460 in sugar, 590 lead chromate, 647, 782 lead compounds, 782 mercury compounds, 783 mineral, 790 non-injurious, 782, 784 orange, 785, 792, 806 orchil, 789 Prussian blue, 790, 792, 810 reagents for identifying, 812 red, 783, 784, 788 references on, 813 Rota's scheme for, 797 separation by solvents, 795 toxic effect of, 781	i *	
Coffee, 379 adulteration of, 384 ash of, 380, 382 caffeine in, 380, 384 caffetannic acid in, 379, 582 caffeol in, 379 chicory in, 388, 389 coloring of, 384 composition of, 379, 380, 381 essential oil of, 379 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384 in sugar, 590 lead chromate, 647, 782 lead compounds, 782 mercury compounds, 783 mineral, 790 non-injurious, 782, 784 orange, 785, 792, 806 orchil, 789 Prussian blue, 790, 792, 810 reagents for identifying, 812 red, 783, 784, 788 references on, 813 Rota's scheme for, 797 separation by solvents, 795 toxic effect of, 781		
ash of, 380, 382 caffeine in, 380, 384 caffetannic acid in, 379, 382 caffeol in, 379 chicory in, 388, 389 coloring of, 384 composition of, 379, 380, 381 essential oil of, 379 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384 lead compounds, 782 logwood, 789 mercury compounds, 783 mineral, 790 non-injurious, 782, 784 orange, 785, 792, 806 orchil, 789 Prussian blue, 790, 792, 810 reagents for identifying, 812 references on, 813 Rota's scheme for, 797 separation by solvents, 795 toxic effect of, 781		
caffeine in, 380, 384 caffetannic acid in, 379, 582 caffeol in, 379 chicory in, 388, 389 coloring of, 384 composition of, 379, 380, 381 essential oil of, 379 fat in, 379 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384 logwood, 789 mercury compounds, 783 mineral, 790 non-injurious, 782, 784 orange, 785, 792, 806 orchil, 789 Prussian blue, 790, 792, 810 reagents for identifying, 812 references on, 813 Rota's scheme for, 797 separation by solvents, 795 toxic effect of, 781	adulteration of, 384	lead chromate, 647, 782
caffetannic acid in, 379, 382 caffeol in, 379 chicory in, 388, 389 coloring of, 384 composition of, 379, 380, 381 essential oil of, 379 fat in, 379 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384 mercury compounds, 783 mineral, 790 non-injurious, 782, 784 orange, 785, 792, 806 orchil, 789 Prussian blue, 790, 792, 810 reagents for identifying, 812 references on, 813 Rota's scheme for, 797 separation by solvents, 795 toxic effect of, 781	ash of, 380, 382	lead compounds, 782
caffeol in, 379 chicory in, 388, 389 coloring of, 384 composition of, 379, 380, 381 essential oil of, 379 fat in, 379 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384 mineral, 790 non-injurious, 782, 784 orange, 785, 792, 806 orchil, 789 Prussian blue, 790, 792, 810 reagents for identifying, 812 references on, 813 Rota's scheme for, 797 separation by solvents, 795 toxic effect of, 781		logwood, 789
chicory in, 388, 389 coloring of, 384 composition of, 379, 380, 381 essential oil of, 379 fat in, 379 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384 non-injurious, 782, 784 orange, 785, 792, 806 orchil, 789 Prussian blue, 790, 792, 810 reagents for identifying, 812 red, 783, 784, 788 references on, 813 Rota's scheme for, 797 separation by solvents, 795 toxic effect of, 781	caffetannic acid in, 379, 382	mercury compounds, 783
coloring of, 384 composition of, 379, 380, 381 essential oil of, 379 fat in, 379 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384 orange, 785, 792, 806 orchil, 789 Prussian blue, 790, 792, 810 reagents for identifying, 812 red, 783, 784, 788 references on, 813 Rota's scheme for, 797 separation by solvents, 795 toxic effect of, 781		mineral, 790
composition of, 379, 380, 381 essential oil of, 379 fat in, 379 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384 orchil, 789 Prussian blue, 790, 792, 810 reagents for identifying, 812 red, 783, 784, 788 references on, 813 Rota's scheme for, 797 separation by solvents, 795 toxic effect of, 781		non-injurious, 782, 784
essential oil of, 379 fat in, 379 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384 Prussian blue, 790, 792, 810 reagents for identifying, 812 red, 783, 784, 788 references on, 813 Rota's scheme for, 797 separation by solvents, 795 toxic effect of, 781		orange, 785, 792, 806
fat in, 379 glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384 reagents for identifying, 812 red, 783, 784, 788 references on, 813 Rota's scheme for, 797 separation by solvents, 795 toxic effect of, 781		
glazing of, 385 hygienic, 390 methods of analysis, 382 microscopical structure, 386 "pellets," 384 red, 783, 784, 788 references on, 813 Rota's scheme for, 797 separation by solvents, 795 toxic effect of, 781		Prussian blue, 790, 792, 810
hygienic, 390 references on, 813 methods of analysis, 382 Rota's scheme for, 797 microscopical structure, 386 separation by solvents, 795 "pellets," 384 toxic effect of, 781		
methods of analysis, 382 microscopical structure, 386 "pellets," 384 Rota's scheme for, 797 separation by solvents, 795 toxic effect of, 781		and the second s
microscopical structure, 386 separation by solvents, 795 toxic effect of, 781		
"pellets," 384 toxic effect of, 781		
	· · · · · · · · · · · · · · · · · · ·	
references on, 400 turmeric, 789		
	references on, 400	turmeric, 789

	la
Colors, ultramarine blue, 791, 810	Cottonseed, 516
vegetable, 787, 789, 795	oil, 516
violet, 784, 786, 810	standards for, 517
wool dyeing, 793, 794	tests for, 517
yellow, 783, 785, 788, 792, 806	stearin, 517
Colostrum, 129	Cotton's cane sugar method 185
Commercial glucose. See Glucose	Coumarin, 853
Compressed yeast, 328	determination, 858
Conalbumin, 262	microscopical structure, 860
Concentrated foods, 257	Crampton and Simon's caramel test, 752
Condensed milk, 186	palm oil tests, 542
as a milk adulterant, 186	Cream, 193
ash of, 189	adulteration of, 195
cane sugar in, 191, 192	cheese, 202
composition of, 187	evaporated, 195
fat in, 189, 191, 192	gelatin in, 195
milk sugar in, 190	methods of analysis, 194
methods of analysis, 188	standards for, 195
proteins in, 190, 192	sucrate of lime in, 196
solids of, 188	test scale, 194
standards for, 188	viscogen in, 196
Confectionery, 645	Cream of tartar, 336
adulteration of, 645	in wine, 702
alcohol in, 649	methods of analysis, 336
arsenic in, 649	Creatin, 46, 211
cane sugar in, 648	Creatinin, 46, 211
colors in, 645, 647	Crême de menthe, 755
dextrin in, 648	Crême de Noyau, 754
glucose in, 648	Crude fiber, 277
invert sugar in, 648	in cereals, 296
lead chromate in, 647	Crustaceans, 256
methods of analysis, 646	Crystals, plant, 90
mineral adulterants, 646	Cucumber pickles, 909
paraffin in, 647	Cudbear, 789
starch in, 648	Cuprammonia, 93
Connective tissue 211,	Curaçoa, 754
Copper salts, 897	Curcuma, 450
determination of, 900, 902	Curcumin, 451
in vinegar, 778	Curd tests in butter, 551, 552, 553
Copra oil, 528	Curing meat, 219
Cordials, 754	Currant (color), 788
analysis of, 755	Curry powder, 450
composition of, 755	Custard powders, 270
Corky tissue, 89	
Corn, 271, 272	Dakota mustard, 460
ash of, 302 .	Date stones, 390
bleaching of canned, 904	Decker-Kunze method for theobromine and
composition of, 271, 272	caffeine, 400
microscopical structure, 309	Defren-O'Sullivan sugar method, 150, 594
oil, 521	Defren's sugar tables, 595
sitosterol in, 522	Desiccated egg, 268
proteins of, 300	Deutyro-albumose, 44, 45
starch, 281	Dextrin, 575
syrup, 575	determination of,
Cornelison's butter color test, 537	in cereals, 295
Corning of meat; 219	in glucose, 632

Marie Control

Dextrin, determination of, in honey, 640	Eggs, references on, 270
in molasses, 624	substitutes for, 269
Dextrose, 573	waterglass as a preservative, 266
determination of, 591, 593, 594,	weights of, 264
598	white of, 262
Diastase, animal, 284	yolk of, 263
in malt extract, 729	Elaidin oil test, 499
starch methods, 283	Elastin, 211
Diabetic foods, 357	Elderberry (color), 788
analyses, 358	Electrolytic apparatus, 608
Dietetics, references on, 49	Elm bark, 428
Distilled liquors, 730	Emergency rations, 257
aldehydes in, 745	Ergot, 313
analytical methods, 745	Erythrodextrin, 575
caramel in, 752	Essential oils, 871, 880
color tests 752, 753	Esters, in distilled liquors, 745
esters in, 745	in imitation flavors, 884
extract in, 745	Ether, ethyl, preparation of absolute, 66
furfural in, 746	petroleum, preparation of, for a sol-
fusel oil in, 746	vent, 66
methyl alcohol in, 749	Eucasin, 158
	Eugenol, 412
opalescence test, 753	
references on, 758	Ewe's milk, 127
Doolittle and Woodruff theine method, 373	Exhausted cloves, 418
Doolittle butter color test, 537	ginger, 450
Double dilution sugar method, 149	tea leaves, 375
Dough, expansion of, 317	vanilla beans, 851
Drains, 17	Exhaust pump, 20
Dry wines, 690	Extraction with immiscible solvents, 68
Dry yeast, 328	volatile solvents, 63
Dubois's salicylic acid method, 826	Extractor, Johnson, 55
Dubosc's saccharimeter, 583	Soxhlet, 63
Dulcin, 845	(n
determination, 846	"Faints," 732
Dupré's color method, 705	Farinaceous infants' foods, 356
color tests 752, 753	Fat globules, 90
Dvorkovitsch theine method, 373	Fat of food, 39
	of meat, 226, 227
Ebulioscope, 675	Fats, edible, 471. See also Oils.
Edam cheese, 202	filtering, 473
Edestan, 44	measuring, 473
Edestin, 299, 300	melting point of, 480
Eggs, 261	methods of analysis, 473
ash of, 264	microscopical examination of, 510
carbohydrates of, 263	paraffin in, 510
cold storage, 267	references on, 561
composition of, 264, 265	weighing, 473
desiccated, 268	Fatty acids, 499
fat of, 264	constants of, 500
frozen, 268	insoluble, 485
lecithin determination, 265	solidifying point of, 500
methods of analysis, 265	soluble, 484
opened, 268	volatile, 841
physical examination of, 207	Fehling processes, 500
physical examination of, 267 preservation of, 266	Fehling processes, 590 gravimetric, 150, 503
preservation of, 266 proteins of, 262	Fehling processes, 590 gravimetric, 150, 593 volumetric, 150, 591

Fehling's solution, 591	Food misbranding, 6
equivalents of, 592	nature and composition of, 39
Fermentation, acetic, 759	official control of, 1
alcoholic, 653	references on, 11
lactic, 129	standards, 4
proteolytic, 158, 202	Fore milk, 128
Fermented liquors, 678	Foreshots, 732
Feser's lactoscope, 163	Formaldehyde, 818
Fibrin, 125	detection of, 180, 820
Fibro vascular tissue, 88	determination of, 181, 819,
Fibroin, 42	821
Filled cheese, 203	in eggs, 268
Fish, analyses of, 255	in milk, 178
preservatives in, 257	Fortified wine, 685, 690
Flavoring extracts, 849	Fresenius' method for colors in pastes, 350
references on, 886	Frozen milk, test for, 129
Flesh foods, 211	meat, 239
references on, 258	Fructose, d-, 574
Fletcher and Allen's tannin method, 371	<i>l-δ-</i> , 574
Floor, 15	Fruit, 274
Flour, 311	candied, 646
absorption test of, 317	composition of, 274
acidity in, 320	essences, artificial, 883, 885
adulteration of, 314	juices, 923, 924
alcohol soluble protein in, 320	methods of proximate analysis, 276
alum in, 315	products, 887
baking tests of, 317	references on, 927
bleaching of, 315	references on, 361
detection 21	sugar. See Levulose.
cold water extract of, 320	sugar-coated, 646
color test of, 317	sugar in, 566
composition of, 312	syrups, 926
damaged, 313	tissues under the microscope, 923
dough test of, 317	Fuchsin, 796
fineness of, 316	Fuel value, 47
gluten in, 319, 322	Funnel, jacketted, 474
inspection, 316	separatory, 67, 68
iodine number of fat of, 320	Furfural, 285
methods of analysis, 316	determination, 746
nitrites in, 321	in distilled liquors, 746
proximate constituents of, 319	in vinegar, 777
salt soluble protein in, 320	Fusel oil, 731
Fluoborates, 835, 836	detection, 746
Fluorides, 835	determination, 747
detection of, 835	Fustic, 788
Fluosilicates, 835, 836	
"Foam" test for butter, 549	Game, composition of, 216
Food adulteration, 5	Gases, in spoiled cans, 891
analysis, commercial, 3	Geerlig's table for dry substances in sugar
from dietetic standpoint, 2	products, 615
general methods, 4	Geissler's carbon dioxide apparatus, 337
references on, 79	Gelatin, 42, 211
concentrated, 257	in cream, 195
economy, references on, 49	in meat, 231
inspection, 3, 6	Gerber's milk centrifuge, 136
references on, 11	Gill and Hatch's oil calorimeter, 495

Gin, 744	Gray's method for water in butter, 532
Ginger, 445	Green colors, 783, 784, 785, 792, 810
adulteration of, 450	Groats, 312
black, 446	Gruyère cheese, 202
cold water extract of, 448	Gums, 89
composition of, 446, 447	Gunning-Arnold nitrogen method, 432
exhausted, 447	Gunning nitrogen methods, 69, 71
extract, standards, 881	Gutzeit arsenic test, 632
liming of, 446	, ,
microscopical structure of, 449	Hæmoglobins, 43
oil of, 446,	Halphen cottonseed oil test, 518
standards, 881	Hanus' iodine absorption method, 491
root, 445	Hefelmann's Bombay mace test, 467
standard, 450	Hehner and Richmond's milk formula, 151
white, 446	Hehner's method for insoluble fatty acids, 486
Gliadin, 42, 298, 299	Heidenhain's tartaric acid method, 340
Globulins, 42, 297	Hemicellulose, 285, 296
Globulose, 44	Hess and Prescott vanillin and coumarin
Glucin, 847	method, 858
Glucose, 575	Hetero-albumose, 44, 45
arsenic in, 632	Hiltner's citral method, 868
composition of, 576	Hilyer's benzoic acid method, 831
d-, 573	Histones, 43
determination of, in honey, 637, 641	Hock wine, 689
in jams and jel-	Hoffmeister's schälchen, 64
lies, 919	Holstein cows, milk from, 162
in molasses, 621	Honey, 633
dextrin in, 632	adulteration of, 636
healthfulness of, 576	American, 634
in beer, 710, 712	analysis of, 639
in butter, 539	Canadian, 634
methods of analysis, 630	composition of, 633, 635, 636
standards for, 576	dextro-rotatory, 635, 636
test for, 632, 641	European, 633
Glucoses, 565	gelatin in, 639
Glutelins, 42	glucose in, 637, 641, 642
Gluten, 298, 299	Hawaiian, 634
Bamihl's test for, 322	invert sugar in, 638, 642
biscuit, 358	methods of analysis, 639
determination of, 319	Honeydew, 636, 642
Gluten flour, 357, 358	Hoods, 16
Glutenin, 42, 298, 300	Hops, 708
Glycerin in vanilla extract, 860	substitutes, 710
in wine, 703	Hordein, 42
Glycerin jelly, 86	Horseflesh, characteristics of, 234
Glycogen, 212	composition of, 222
detection, 235	detection of, 235, 237
determination, 236	glycogen in, 235
Glycoproteins, 43	Horseradish, 910
Goat's milk, 127 Gooch's boric acid method, 824	Hortvet method for acids in wine, 701 number, of maple products, 628
Graham flour, 312	of vinegar, 768
Grain, moisture in, 278 Grape juice, 924	and West's benzaldehyde method,
Grape sugar. See Dextrose	875
Grape sugar, standard, 574	rose oil method, 883 spice oil method, 882
Stupe sugar, standard, 3/4	spice on inclide, 662

Hortvet and West's wintergreen oil method, 878	Invert sugar, 589 detection of, 589, 625, 642
Howard's test for gums in ice cream, 201	determination of, 589, 598
volatile oil method, 865	in honey, 638, 642
Hübl's iodine absorption method, 487	Iodine absorption of oils, 487, 491, 492
Human milk, 127	Iodine in potassium iodide, 91
Hungarian red pepper, 439, 441, 442	Irish whiskey, 732, 734, 735
Hunt's iodine reagent, 492	T
Hydrocyanic acid, 874, 877	Jams, 910
Hydrometer, 55	adulteration of, 911, 915
Hypoxanthin, 211	agar agar in, 922
	apple stock in, 914
Ice cream, 198	coagulator in, 914
analytical methods, 199	coloring matter in, 921
detection of thickeners, 200	composition of, g11, g13, g16
fat in, 199	dextrin in, 920
gelatine in, 201	gelatin in, 922
preservatives in, 201	glucose in, 919
standards, 199	methods of analysis, 916
starch in, 201	polarization of, 917
Imitation coffee, 384	preservatives in, 922
Immersion refractometer, 100, 111	starch in, 922
adjustment of scale, 113	sugars in, 917-919
distilled water readings on, 113	Jellies, see Jams
investigation of small quantities of	Johnson extractor, 65
solutions by, 115	Juckenack's lecithin phosphoric acid method,
of solutions excluded from air	, ·
_	349
by, 115	Konwiele's terrerie said method are
milk examination by, 166	Kenrick's tartaric acid method, 340
scale readings compared with n_{D_i}	Kephir, 159
II6	Keratins, 42
solutions standardized by, 120	Ketchups, 905
references on, 122	colors in, 907
temperature corrections for, 121	preservatives in, 908
Incinerator, 173	Kjeldahl nitrogen method, 72
Indicators, 38	Knorr's carbon dioxide apparatus, 338
Indices of refraction, 105, 116	Koelner's baking test, 317
Indigo, 790, 811	Koettstorfer's saponification method, 486
Indol, 92	König and Karach's method for distinguish-
Infants' foods, 354	ing honeydew and glucose, 642
classification of, 355	Koumis, 158
cold water extract of, 360	Kröber's table for pentosans and pentoses,
composition of, 356	288
methods of analysis, 359	
microscopical examination of,	Laboratory benches, 15
360	stain for, 16
preparation of, 355	drains, 17
Inosite, 276	equipment, 14, 15
Inspection of foods, 3, 5, 6, 9	references on, 38
flour, 316	floor, 15
liquors, 655	hoods, 16
milk, 159	lighting, 15
Inulin, 276	location, 14
Invalids' foods, 354. See also Infants'	sinks, 17
Foods.	
_	ventilation, 15
Inversion, 588	Lactalbumin, 125

Lactated infants' foods, 356	Legumes, 272
Lactoglobulin, 125	ash of, 302
Lactometer, 131	Legumin, 42, 300
Lactoscope, 163	Lemon extract, 861
Lactose, 125, 577	adulteration of, 862
Defren's table for, 595	alcohol in, 866
detection of, 625	citral in, 866
determination of, 593, 594, 998, 626	citric acid in, 870
in milk, 126, 127, 147	colors in, 869
Munson and Walker's table for,	composition of, 863
-	lemon oil in, 863, 864
599 Soublet's table for the	methods of analysis, 863
Soxhlet's table for, 152	methyl alcohol in, 869
Lager beer, 708	standard for, 861
Lamb, composition of, 215	tartaric acid in, 870
cuts of, 215	terpeneless, 862
Landwehr's glycogen method, 236	
Lard, 554	oil, terpeneless, 862
adulteration of, 556	Lemongrass oil, 863, 872
back, 554	Lemon juice, 925
composition of, 554	Lemon oil 861, 871
composition of as effected by teed-	determination of, 863, 864
ing, 560	examination of, 870
"compound," 556	Lentils, 272
constants of, 555	Leucosin, 41, 299, 300
iodine number, 559	Levallois' bromine absorption method,
kettle rendered, 554	49.3
leaf, 554	Levulose, 574
microscopical examination of, 557	determination of, 626, 640
neutral, 554	Liebig's meat extract, 242
oil, 555	Lighting, 15
references on, 563	Lignin, 94
standards, 556	Lime, determination of, 303
stearin, 555	in baking powder, 345
substitutes, 559	in spices, 410
Laurent's saccharimeter, 583	juice, 924
La Wall and Bradshaw benzoic acid method,	sucrate of, 196
830	water, in vinegar analysis, 765
Leach and Lythgoe method for malic value	Liming of ginger, 446
in maple products, 627	Limonene, 871
methyl alcohol method, 749	Liqueurs, 754
Lead chromate, 647, 782	analysis of, 755
number, maple products, 628	Liquor inspection, 655
vinegar, 768	Liquors, alcohol in, 658, 715
salts of, 892, 896	ash of, 677
determination of, 899, 900, 902	distilled, 730
Leavening materials, 327.332	methods of analysis, 745
references on, 364	extract of, 677
Lecitalbumin, 13	fermented, 678
Lecithin, 46	malt, 707
determination of, 265	methods of analysis, 714
nucleovitellin, 43	malted and non-malted, 712 -
Lecithoproteins, 43	methods of analysis, 657
Leffmann and Beam's method for volatile	preservatives in, 677
fatty acids, 482	specific gravity of, 657
fat method, 49	Lobster, composition of, 256
Legumelin, 41	Logwood, 789
J , ,	U / 1 - 7

	1
Long fermentation baking test, 318	Meat, antiseptics in, 220
pepper, 438	ash in, 225
Lovibond tintometer, 77	bases, 211, 222, 228, 231
Lowenthal's tannin method, 370	boric acid in, 232
Low wines, 732	canned, 221
Lythgoe's sucrose test for milk, 197	canning of, 221
=,g, -,,	colors in, 238
Macaroni, 347	composition of, 221
Macassar mace, 468	cooking, effect of, 220
Mace, 462, 465	corning of, 219
adulteration of, 466	curing of, 219
	extracts, 240
Bombay, 467	
composition of, 465	acidity of, 253
Macassar, 468	albumoses in, 250
microscopical structure of, 466	ash in, 249
standard, 466	composition of, 242, 243,
Madeira wine, 687	247
Maize. See Corn.	creatin in, 244, 252
Malaga wine, artificial, 692	creatinin in, 244, 252
Malic acid in cider, 702	fat in, 249
in vinegar, 767	fluid, 241, 243, 244
in wine, 702	gelatin in, 253
value in maple products, 627	glycerol in, 254
Malt, 707	meat bases in, 252
extracts, 284, 729	methods of analysis, 246
liquors, 707. See also Beer.	nitrogen compounds of, 249,
substitutes, 710	250
vinegar, 762	peptones in, 251
Malting, 707	preservatives in, 254
Maltose, 574	proteoses in, 250
detection of, 625	solid, 241, 242, 244
determination of, 594, 598 626	standards, 241
Maple sap, 570	xanthin bases in, 253
sugar, 570. See also Maple syrup.	fat, composition of, 226
syrup, 570	determination, 226
adulteration of, 572	gelatin determination, 231
ash of, 571, 572	glycogen in, 236
composition of, 571, 572	inspection, 217
Hortvet number of, 628	juices, 241, 245, 247, 248
lead number of, 628	manufactured, 218
malic acid value, 627	methods of analysis, 225
methods of analysis, 627	nitrates in, 232
moisture in, 627	nitrogen determination, 226
standards, 572	nitrogenous bodies, separation of, 228
Maraschino, 754	peptones in, 251
Mare's milk, 127	pickled, 218
Marigold, 789	powders, 247, 248
Marpmann's color method, 239	preservation of, 218
Marsh arsenic test, 75, 728	preservatives in, 232
test for caramel, 753	proteins, coagulable, 231
Martin's color scheme, 535	proteoses in, 231
"Materna" milk modifier, 157	ptomaines in, 218
Maumené thermal test, 494	refrigeration of, 210
Mayrhofer's glycogen method, 237	salicylic acid in, 233
McGill's drying oven, 586	salted, 219
Meat, 211	smoked, 210
,	

Meat, standards of, 218 sulphurous acid in, 231 unwholesome, 218 water in, 225 Melting point, 480 Mercury compounds in colors, 783 Metallic salts in canned goods, toxic effects of, 899 Metaproteins, 44 Micro-polariscope, 84 Microscopi in food analysis, 81 references on, 98 reagents for, 90 stand, 82 Microscopical accessories, 84 analysis, 81 apparatus, 82 diagnosis, 86 reagents, 90 starch, 281 bean, 388 starch, 282 buckwheat, 311, 437 starch, 281 butter, 552 cassia 426 cayenne, 441 cereal products, 305 chicory, 386 cinnamon, 426 cloves 416 cocoa, 403 cocoanut shells, 419 coffee, 386 corn, 390 starch, 281 date stones, 390 fats, 510 flour, 306, 322 ginger, 449 honey, 633 jams, 923 jellies, 943 lard, 557 mace, 466 milk, 124 mustard, 458 nutmeg, 464 Microscopy of oats, 390 oat starch, 282 oils, 510 oleomargarine, 552 oilevate, 522 olive stones, 436 paprika, 441 pea, 388 starch, 282 pepper, black, 433 long, 439 red, 441 white, 433 potato starch, 282 ree, 308 starch, 282 ree, 308 starch, 282 ree, 308 starch, 282 ree, 308 starch, 282 pepper, black, 433 long, 439 red, 441 white, 433 say, 283 sawdust, 444 starches, 280 starch, 282 ree, 308 starch, 282 pepper, black, 433 long, 439 red, 441 white, 433 say, 283 sawdust, 444 starches, 280 starch, 282 ree, 308 starch, 282 pepper, black, 433 long, 439 red, 441 white, 433 say, 283 sawdust, 444 starches, 280 starch, 281 sago, 283 sawdust, 444 starches, 280 starch, 281 sago, 283 sawdust, 444 starches, 280 starch, 281 sago, 283 sawdust, 444 starches, 280 starch, 281 sago, 283 sawdust, 444 starches, 280 starch, 281 sago, 283 sawdust, 444 starches, 280 starch, 281 sago, 283 sawdust, 444 starches, 280 starch, 281 sago, 283 sawdust, 444 starches, 280 starch, 281 sago, 283 sawdust, 444 starches, 280 starch, 281 sago, 283 sawdust, 444 starches, 280 starch, 281 sago, 283 sawdust, 444 starches, 280 starch, 281 sago, 283 sawdust, 444 starches, 280 starch, 281 sago, 283 sawdust, 444 starches, 280 starch, 281 sago, 283 sawdust, 444 starches, 280 starch, 281 sago, 283 sawdust, 444 starches,		
unwholesome, 218 water in, 225 Melting point, 480 Mercury compounds in colors, 783 Metallic salts in canned goods, toxic effects of, 899 Metaproteins, 44 Methyl alcohol, detection of, 749, 869 Micro-chemical reactions, 94 Micro-polariscope, 84 Microscope in food analysis, 81 references on, 98 reagents for, 90 stand, 82 Microscopical accessories, 84 analysis, 81 apparatus, 82 diagnosis, 86 reagents, 90 analytical, 91 clarifying, 92 Microscopy of agar agar, 922 allspice, 422 arrowroot, 282 barley, 309 starch, 281 bean, 388 starch, 282 buckwheat, 311, 437 starch, 281 butter, 552 cassia 426 cayenne, 441 cereal products, 305 chicory, 386 cinnamon, 426 cloves 416 cocoa, 403 cocoanut shells, 419 coffee, 386 corn, 309 starch, 281 date stones, 390 fats, 510 flour, 306, 322 ginger, 449 honey, 633 jams, 923 jellies, 923 lard, 557 mace, 466 milk, 124 mustard, 458 olicomargarine, 552 olice stones, 436 paprika, 441 pea, 388 starch, 282 pepper, black, 433 long, 439 red, 441 white, 433 paporatos starch, 282 rice, 310 starch, 282 rice, 310 starch, 282 starch, 281 sago, 283 sawdust, 444 starches, 280 tarch, 281 sago, 283 turneric, 451 wheat, 311, 437 starch, 281 sago, 283 turneric, 451 wheat, 313 wheat, 343 bons, 439 red, 441 white, 433 paporatos starch, 282 rice, 310 starch, 282 rice, 310 starch, 282 starch, 282 rice, 310 starch, 282 rice, 310 starch, 282 starch, 282 rice, 310 starch, 282 starch, 282 starch, 282 rice, 310 starch, 282 starch, 282 starch, 282 sago, 283 sawdust, 444 starches, 280 starch, 282 starch, 281 sago, 283 sawdust, 444 starches, 280 starch, 281 sago, 283 sawdust, 444 starches, 280 starch, 281 sago, 283 sawdust, 444 starches, 280 starch, 281 sago, 283 sawdust, 444 starches, 280 starch, 281 sago, 283 sawdust, 444 starches, 280 starch, 281 sago, 283 starch, 281 sago, 283 starch, 281 sago, 283 starch, 281 sago, 283 starch, 281 sago, 283 starch, 281 sago, 283 starch, 281 sago, 283 sawdust, 444 starches, 280 starch, 281 sago, 283 starch, 281 sago, 283 sawdust, 444 starches, 280 starch, 281 sago, 283 sawdust, 4		
water in, 225 Melting point, 480 Mercury compounds in colors, 783 Metallic salts in canned goods, toxic effects of, 899 Metaproteins, 44 Micro-tenins, 44 Micro-polariscope, 84 Microscope in food analysis, 81		
Melting point, 480 Mercury compounds in colors, 783 Meratury compounds in colors, 783 Metallic salts in canned goods, toxic effects of, 899 Metaproteins, 44 Methyl alcohol, detection of, 749, 869 Micro-chemical reactions, 94 Micro-polariscope, 84 Microscope in food analysis, 81 references on, 98 reagents for, 90 stand, 82 Microscopical accessories, 84 analysis, 81 apparatus, 82 diagnosis, 86 reagents, 90 analytical, 91 clarifying, 92 Microscopy of agar agar, 922 allspice, 422 arrowroot, 282 barley, 309 starch, 281 bean, 388 starch, 281 bean, 388 starch, 281 buckwheat, 311, 437 starch, 281 butter, 552 cassia 426 cayenne, 441 cereal products, 305 chicory, 386 cinnamon, 426 cloves 416 cocoa, 403 cocoanut shells, 419 coffee, 386 corn, 309 starch, 281 date stones, 390 fats, 510 flour, 306, 322 ginger, 449 honey, 633 jams, 923 jellies, 923 lard, 557 mace, 466 milk, 124 mustard, 458 olive stones, 436 paprika, 441 pea, 388 starch, 282 pepper, black, 433 long, 439 red, 441 white, 433 potato starch, 282 rice, 310 starch, 282 starch, 282 rice, 310 starch, 282 rice, 310 starch, 282 starch, 282 rice, 310 starch, 282 starch, 282 rice, 310 starch, 282 starch, 282 rice, 310 starch, 282 starch, 282 rice, 310 starch, 282 starch, 281 starch, 281 starch, 281 starch, 281 starch, 281 starch, 281 starch, 281 starch, 281 starch	•	
Mercury compounds in colors, 783 Metallic salts in canned goods, toxic effects of, 899 Metaproteins, 44 Methyl alcohol, detection of, 749, 869 Micro-chemical reactions, 94 Micro-chemical reactions, 94 Micro-chemical reactions, 94 Microscope in food analysis, 81 references on, 98 reagents for, 90 stand, 82 Microscopical accessories, 84 analysis, 81 apparatus, 82 diagnosis, 86 reagents, 90 analytical, 91 clarifying, 92 Microscopy of agar agar, 922 allspice, 422 arrowroot, 282 barley, 309 Starch, 281 bean, 388 starch, 282 buckwheat, 311, 437 starch, 281 bean, 388 starch, 282 buckwheat, 311, 437 starch, 281 bean, 388 starch, 282 diagnosis, 86 cayenne, 441 cereal products, 305 chicory, 386 cinnamon, 426 cloves 416 cocoa, 403 cocoanut shells, 419 coffee, 386 corn, 309 starch, 281 date stones, 390 fats, 510 flour, 306, 322 ginger, 449 honey, 633 jams, 943 jellies, 923 lard, 557 mace, 466 milk, 124 mustard, 458		
Metalic salts in canned goods, toxic effects of, 899 Metaproteins, 44 Methyl alcohol, detection of, 749, 869 Micro-chemical reactions, 94 Micro-polariscope, 84 Microscope in food analysis, 81		
of, 899 Metaproteins, 44 Methyl alcohol, detection of, 749, 869 Micro-chemical reactions, 94 Micro-polariscope, 84 Microscope in food analysis, 81 references on, 98 reagents for, 90 stand, 82 Microscopical accessories, 84 analysis, 81 apparatus, 82 diagnosis, 86 reagents, 90 analytical, 91 clarifying, 92 Microscopy of agar agar, 922 allspice, 422 arrowroot, 282 barley, 399 starch, 281 bean, 388 starch, 281 beuckwheat, 311, 437 starch, 281 butter, 552 cassia 426 cayenne, 441 cereal products, 305 chicory, 386 cinnamon, 426 cloves 416 cocoa, 403 cocoanut shells, 419 coffee. 386 corn, 399 starch, 281 date stones, 390 fats, 510 flour, 306, 322 ginger, 449 honey, 633 jams, 923 jellies, 923 lard, 557 mace, 466 milk, 124 mustard, 458 starch, 282 pepper, black, 433 long, 439 red, 441 white, 433 potato starch, 282 rice, 310 starch, 282 rice, 310 starch, 282 rice, 310 starch, 282 rice, 310 starch, 282 rice, 310 starch, 282 starch, 282 rice, 310 starch, 282 rice, 310 starch, 282 rice, 310 starch, 282 rice, 310 starch, 282 rice, 310 starch, 282 rice, 310 starch, 282 rice, 310 starch, 282 starch, 282 rice, 310 starch, 282 rice, 310 starch, 282 rice, 310 starch, 282 rice, 310 starch, 282 starch, 282 starch, 281 sago, 283 audustration of, 155 soric acid in, 124 ass's, 127 hoiled milk, detection, 155 boric acid in, 182 calcium oxide in, 188 calculation of proteins, 153 caramel in, 176, 177 carbonate in, 180, 182 chocolate, 397 citric acid in, 127 coloring matter in, 174–177 composition of, 124–126 constants, 169 ewe's, 127 fat of, 127, 134 fermentations of, 129 foods, prepared, 157 fore milk, 128 formaldebyde in, 178, 181 goat's, 117 human, 127 inspection, 159		
Metaproteins, 44 Methyl alcohol, detection of, 749, 869 Micro-chemical reactions, 94 Micro-polariscope, 84 Microscope in food analysis, 81	Metallic salts in canned goods, toxic effects	
Methyl alcohol, detection of, 749, 869 Micro-chemical reactions, 94 Micro-polariscope, 84 Microscope in food analysis, 81	` . * *	
Micro-chemical reactions, 94 Micro-polariscope, 84 Microscope in food analysis, 81		
Micro-polariscope, 84 Microscope in food analysis, 81	•	
Microscope in food analysis, 81	Micro-chemical reactions, 94	
references on, 98 reagents for, 90 stand, 82 Microscopical accessories, 84 analysis, 81 apparatus, 82 diagnosis, 86 reagents, 90 analytical, 91 clarifying, 92 Microscopy of agar agar, 922 allspice, 422 arrowroot, 282 barley, 309 starch, 281 bean, 388 starch, 282 buckwheat, 311, 437 starch, 281 butter, 552 cassia 426 cayenne, 441 cereal products, 305 chicory, 386 cinnamon, 426 cloves 416 cocoa, 403 cocoanut shells, 419 coffee, 386 corn, 309 starch, 281 date stones, 390 fats, 510 flour, 306, 322 ginger, 449 honey, 633 jams, 923 jellies, 923 lard, 557 mace, 466 milk, 124 mustard, 458 microscopical accessories, 84 analysis, 81 sago, 282 starch, 281 sago, 283 sawdust, 444 starches, 280 tapioca starch, 282 Microscopical accessories, 84 saporatus, 82 starch, 281 sago, 283 sawdust, 444 starches, 280 tapioca starch, 282 Mik, 124 acidity of, 124, 153 adulteration of, 159 alkalinity of ash, 198 anilin orange in, 175, 177 annatto in, 175, 177 annatto in, 175, 177 annatto in, 175, 177 annatto in, 175, 177 annatto in, 175, 177 carbonate in, 180, 182 chocolate, 397 citric acid in, 127 coloring matter in, 174-177 composition of, 124-126 constants, 169 ewe's, 127 fat of, 127, 134 fermentations of, 129 foods, prepared, 157 fore milk, 128 formaldehyde in, 178, 181 goat's, 127 human, 127 inspection, 159		
reagents for, 90 stand, 82 Microscopical accessories, 84 analysis, 81 apparatus, 82 diagnosis, 86 reagents, 90 analytical, 91 clarifying, 92 Microscopy of agar agar, 922 allspice, 422 arrowroot, 282 barley, 309 starch, 281 bean, 388 starch, 282 buckwheat, 311, 437 starch, 281 butter, 552 cassia 426 cayenne, 441 cereal products, 305 chicory, 386 cinnamon, 426 cloves 416 cocoa, 403 cocoanut shells, 419 coffee. 386 corn, 309 starch, 281 date stones, 390 fats, 510 flour, 306, 322 ginger, 449 honey, 633 jams, 923 jellies, 923 lard, 557 mace, 466 milk, 124 mustard, 458 Microscopy of sarch, 281 sago, 283 sawdust, 444 starches, 280 tapioca starch, 282 thea, 378 turmeric, 451 wheat, 3c6 starch, 281 sadoust, 424 acidity of, 124, 153 adulteration of, 159 alkalinity of ash, 198 anilin orange in, 175, 177 annatto in, 175, 176 ash of, 127, 134 ashing of, 134 ass's, 127 boiled milk, detection, 155 boric acid in, 182 calcium oxide in, 198 calculation of proteins, 153 caramel in, 176, 177 composition of, 124-126 constants, 169 ewe's, 127 fat of, 127, 134 fermentations of, 129 foods, prepared, 157 fore milk, 128 formaldehyde in, 178, 181 goat's, 127 human, 127 inspection, 159		
stand, 82 Microscopical accessories, 84	_ , , ,	
Microscopical accessories, 84	reagents for, 90	starch, 282
analysis, 81 apparatus, 82 diagnosis, 86 reagents, 90 analytical, 91 clarifying, 92 Microscopy of agar agar, 922 arrowroot, 282 barley, 309 starch, 281 bean, 388 starch, 282 buckwheat, 311, 437 starch, 281 butter, 552 cassia 426 cayenne, 441 cereal products, 305 chicory, 386 cinnamon, 426 cloves 416 cocoa, 403 cocoanut shells, 419 coffee. 386 corn, 309 starch, 281 date stones, 390 fats, 510 flour, 306, 322 ginger, 449 honey, 633 jams, 923 jellies, 923 lard, 557 mace, 466 milk, 124 mustard, 458 Microscopy of tagar agar, 922 arrowroot, 282 buckwheat, 378 turmeric, 451 wheat, 378 turmeric, 451 wheat, 376 date, 378 turmeric, 451 wheat, 376 date, 376 diamico-technique, 82 Milk, 124 acidity of, 124, 153 adulteration of, 159 alkalinity of ash, 198 anilin orange in, 175, 177 annatto in, 175, 177 ash of, 127, 134 ashing of, 13	stand, 82	rye, 308
apparatus, 82 diagnosis, 86 reagents, 90 analytical, 91 clarifying, 92 Microscopy of agar agar, 922 allspice, 422 arrowroot, 282 barley, 309 starch, 281 bean, 388 starch, 282 buckwheat, 311, 437 starch, 281 butter, 552 cassia 426 cayenne, 441 cereal products, 305 chicory, 386 cinnamon, 426 cloves 416 cocoa, 403 cocoanut shells, 419 coffee, 386 corn, 309 starch, 281 date stones, 390 fats, 510 flour, 366, 322 ginger, 449 honey, 633 jams, 923 jellies, 923 lard, 557 mace, 466 milk, 124 mustard, 458 sawdust, 444 starches, 280 tapioca starch, 282 ttea, 378 turmeric, 451 wheat, 3c6 starch, 281 Micro-technique, 82 acidity of, 124, 153 adulteration of, 159 anhitt of ash, 198 anilin orange in, 175, 177 annatto in, 175, 176 ash of, 127, 134 assi's, 127 hoiled milk, detection, 155 boric acid in, 182 calcium oxide in, 198 calculation of proteins, 153 caramel in, 176, 177 composition of, 124–126 constants, 169 ewe's, 127 fat of, 127, 134 fermentations of, 129 foods, prepared, 157 fore milk, 128 formaldehyde in, 178, 181 goat's, 127 human, 127 inspection, 159		starch, 281
diagnosis, 86 reagents, 90	analysis, 81	
reagents, 90		sawdust, 444
analytical, 91 clarifying, 92 Microscopy of agar agar, 922 arrowroot, 282 barley, 309 starch, 281 bean, 388 starch, 282 buckwheat, 311, 437 starch, 281 butter, 552 cassia 426 cayenne, 441 cereal products, 305 chicory, 386 cinnamon, 426 cloves 416 cocoa, 403 cocoanut shells, 419 coffee. 386 corn, 309 starch, 281 date stones, 390 fats, 510 flour, 306, 322 ginger, 449 honey, 633 jams, 923 jellies, 923 lard, 557 mace, 466 milk, 124 mustard, 458 Micro-technique, 82 Mik, 124 acidity of, 124, 153 adulteration of, 159 alkalinity of ash, 198 anilin orange in, 175, 177 annatto in, 175, 176 ash of, 127, 134 ashing of, 134 assi's, 127 hoiled milk, detection, 155 boric acid in, 182 calcium oxide in, 188 calculation of proteins, 153 caramel in, 176, 177 carbonate in, 180, 182 chocolate, 397 citric acid in, 127 coloring matter in, 174–177 composition of, 124–126 constants, 169 ewe's, 127 fat of, 127, 134 fermentations of, 129 foods, prepared, 157 fore milk, 128 mustard, 458	diagnosis, 86	starches, 280
Clarifying, 92 Clar	reagents, 90	tapioca starch, 282
Microscopy of agar agar, 922 allspice, 422 arrowroot, 282 barley, 309 starch, 281 bean, 388 starch, 282 buckwheat, 311, 437 starch, 281 butter, 552 cassia 426 cayenne, 441 cereal products, 305 chicory, 386 cinnamon, 426 cloves 416 cocoa, 403 cocoanut shells, 419 coffee, 386 corn, 309 starch, 281 date stones, 390 fats, 510 flour, 306, 322 ginger, 449 honey, 633 jams, 923 jellies, 923 lard, 557 mace, 466 milk, 124 mustard, 458 Micro-technique, 82 Acidity of, 124, 153 adulteration of, 159 alkalinity of ash, 198 anilin orange in, 175, 177 annatto in, 175, 170 ash of, 127, 134 ass's, 127 boiled milk, detection, 155 boric acid in, 182 calcium oxide in, 198 calculation of proteins, 153 caramel in, 176, 177 carbonate in, 180, 182 chocolate, 397 citric acid in, 127 coordinate in, 180, 182 chocolate, 397 citric acid in, 127 coordinate in, 180, 182 chocolate, 397 coordinate in, 174-177 corbonate in, 180, 182 chocolate, 397 coordinate in, 174-177 corbonate in, 127 inspection, 129 foods, prepared, 157	analytical, 91	tea, 378
allspice, 422 arrowroot, 282 barley, 309 starch, 281 bean, 388 starch, 282 buckwheat, 311, 437 starch, 281 butter, 552 cassia 426 cayenne, 441 cereal products, 305 chicory, 386 cinnamon, 426 cloves 416 cocoa, 403 coroanut shells, 419 coffee. 386 corn, 309 starch, 281 date stones, 390 fats, 510 flour, 306, 322 ginger, 449 honey, 633 jams, 923 jellies, 923 jellies, 923 jellies, 923 jellies, 923 jellies, 923 jams, 923 jellies, 923 jellies, 923 jellies, 923 jellies, 923 jellies, 923 jellies, 923 jellies, 946 milk, 124 mustard, 458 Micro-technique, 82 Milk, 124 microtethnique, 82 Milk, 124 microtechnique, 82 Milk, 124 microtechnique, 82 Milk, 124 microtechnique, 82 Milk, 124 microtechnique, 82 Milk, 124 milk, 124 milk, 124 milk, 124 milk, 124 milk, 124 milk, 124 milk, 124 milicrotechnique, 82 Milk, 124 milk, 124 milk, 124 milcrotechnique, 82 Milk, 124 milk	clarifying, 92	turmeric, 451
arrowroot, 282 barley, 309	Microscopy of agar agar, 922	
barley, 309	allspice, 422	starch, 281
starch, 281 bean, 388 starch, 282 buckwheat, 311, 437 starch, 281 butter, 552 cassia 426 cayenne, 441 cereal products, 305 chicory, 386 cinnamon, 426 cloves 416 cocoa, 403 corn, 309 starch, 281 coffee, 386 corn, 309 starch, 281 date stones, 390 fats, 510 flour, 306, 322 ginger, 449 honey, 633 jams, 923 jellies, 923 lard, 557 mace, 466 milk, 124 mustard, 458 acidity of, 124, 153 adulteration of, 159 alkalinity of ash, 198 anilin orange in, 175, 177 annatto in, 175, 177 annatto in, 175, 176 ash of, 127, 134 ass's, 127 boiled milk, detection, 155 boric acid in, 182 calcium oxide in, 198 calculation of proteins, 153 caramel in, 176, 177 carbonate in, 180, 182 chocolate, 397 citric acid in, 127 constants, 169 ewe's, 127 fat of, 124, 153 adulteration of, 159 alkalinity of ash, 198 anilin orange in, 175, 177 annatto in, 175, 177 ash of, 127, 134 ass's, 127 boiled milk, detection, 155 boric acid in, 182 calcium oxide in, 198 calculation of, 159 alkalinity of ash, 198 anilin orange in, 175, 177 annatto in, 175, 177 ash of, 127, 134 ass's, 127 boiled milk, detection, 155 boric acid in, 182 calcium oxide in, 198 calculation of, 159 alkalinity of ash, 198 anilin orange in, 175, 177 annatto in, 175, 176 ash of, 127, 134 ass's, 127 boiled milk, detection, 155 boric acid in, 182 calcium oxide in, 198 calculation of, 124 ass's, 127 boiled milk, detection, 155 boric acid in, 182 calcium oxide in, 198 calculation of, 124 ass's, 127 annatto in, 175, 176 ash of, 127, 134 ass's, 127 colering milk, 124 ass's, 127 annatto in, 175, 176 ash of, 127, 134 ass's, 127 colering milk, 124 ass's, 127 annatto in, 175, 176 ash of, 127, 134 ass's, 127 calcium oxide in, 198 calcium oxide in, 198 calcium oxide in, 198 calcium oxide in, 198 calcium oxide in, 198 calcium oxide in, 198 calcium oxide in, 198 calcium oxide in, 198 calcium oxide in, 198 calcium oxide in, 198 calcium oxide in, 198 calcium oxide in, 198 calcium oxide in, 198 calcium oxide in, 198 calcium oxide in, 198 calcium oxide in, 198 calcium oxide in, 198 calcium oxide in, 198 calc	arrowroot, 282	Micro-technique, 82
bean, 388		Milk, 124
starch, 282 buckwheat, 311, 437 starch, 281 butter, 552 cassia 426 cayenne, 441 cereal products, 305 chicory, 386 cinnamon, 426 cloves 416 cocoa, 403 corn, 309 starch, 281 date stones, 390 fats, 510 flour, 306, 322 ginger, 449 honey, 633 jams, 923 jellies, 923 lard, 557 mace, 466 milk, 124 mustard, 458 alkalinity of ash, 198 anilin orange in, 175, 177 annatto in, 175, 176 ash of, 127, 134 assi's, 127 annatto in, 175, 176 ash of, 127, 134 assi's, 127 annatto in, 175, 176 ash of, 127, 134 assi's, 127 calcium oxide in, 198 calculation of proteins, 153 caramel in, 176, 177 carbonate in, 180, 182 chocolate, 397 citric acid in, 127 coloring matter in, 174–177 composition of, 124–126 constants, 169 ewe's, 127 fat of, 127, 134 fermentations of, 129 foods, prepared, 157 fore milk, 128 formaldehyde in, 178, 181 goat's, 127 human, 127 inspection, 159	•	
buckwheat, 311, 437	hean, 388	
starch, 281 butter, 552 cassia 426 cayenne, 441 cereal products, 305 chicory, 386 cinnamon, 426 cloves 416 cocoa, 403 corn, 309 starch, 281 date stones, 390 fats, 510 flour, 306, 322 ginger, 449 honey, 633 jams, 923 jams, 923 jellies, 923 lard, 557 mace, 466 milk, 124 mustard, 458 annatto in, 175, 176 ash of, 127, 134 ashing of, 134 ass's, 127 hoiled milk, detection, 155 boric acid in, 182 calcium oxide in, 198 calculation of proteins, 153 caramel in, 176, 177 carbonate in, 180, 182 chocolate, 397 citric acid in, 127 constants, 169 ewe's, 127 fat of, 127, 134 fermentations of, 129 foods, prepared, 157 fore milk, 128 formaldehyde in, 178, 181 mystard, 458	starch, 282	
butter, 552 cassia 426 cayenne, 441 cereal products, 305 chicory, 386 cinnamon, 426 cloves 416 cocoa, 403 corn, 309 starch, 281 date stones, 390 fats, 510 flour, 306, 322 ginger, 449 honey, 633 jams, 923 jams, 923 jams, 923 jard, 557 mace, 466 milk, 124 mustard, 458 ash of, 127, 134 assing of, 134 assi's, 127 holied milk, detection, 155 boric acid in, 182 calcium oxide in, 198 calcium oxide i		anilin orange in, 175, 177
cassia 426 cayenne, 441 cereal products, 305 chicory, 386 cinnamon, 426 cloves 416 cocoa, 403 corr, 309 corr, 309 starch, 281 date stones, 390 fats, 510 flour, 306, 322 ginger, 449 honey, 633 jams, 923 jams, 923 jams, 923 jard, 557 mace, 466 milk, 124 mustard, 458 ashing of, 134 ass's, 127 hoiled milk, detection, 155 boric acid in, 182 calcium oxide in, 198 calcium oxide	starch, 281	annatto in, 175, 176
cayenne, 441 cereal products, 305 chicory, 386 cinnamon, 426 cloves 416 cocoa, 403 corfiee. 386 corn, 309 starch, 281 date stones, 390 fats, 510 flour, 306, 322 ginger, 449 honey, 633 jams, 923 jellies, 923 lard, 557 mace, 466 milk, 124 mustard, 458 ass's, 127 hoiled milk, detection, 155 boric acid in, 182 calcium oxide in, 198 calculation of proteins, 153 caramel in, 176, 177 carbonate in, 180, 182 chocolate, 397 citric acid in, 127 coloring matter in, 174-177 composition of, 124-126 constants, 169 ewe's, 127 fat of, 127, 134 fermentations of, 129 foods, prepared, 157 fore milk, 128 formaldehyde in, 178, 181 goat's, 127 human, 127 inspection, 159		
cereal products, 305 chicory, 386 cinnamon, 426 cloves 416 cocoa, 403 cocoanut shells, 419 coffee. 386 corn, 309 starch, 281 date stones, 390 fats, 510 flour, 306, 322 ginger, 449 honey, 633 jams, 923 jams, 923 jams, 923 jars, 923 lard, 557 mace, 466 milk, 124 mustard, 458 hoiled milk, detection, 155 boric acid in, 182 calcium oxide in, 198 cakulation of proteins, 153 caramel in, 176, 177 carbonate in, 180, 182 cabonate in, 180, 182 carbonate in, 180, 182 cabonate in, 180, 182 carbonate in, 180, 182 carbonate in, 180, 182 carbonate in, 180, 182 cabonate in, 180, 182 cabonate in, 198 carbonate in, 180, 182 cabonate in, 198 carbonate in, 180, 182 cabonate in, 198 carbonate in, 180, 182 cabonate in, 180, 182 cabonate in, 198 carbonate in, 198 carbonate in, 180, 182 cabonate in, 180, 182 cabonate in, 180 carbonate in, 198 carbonate in, 198 carbonate in, 198 carbonate in, 198 carbonate in, 198 carbonate in, 198 carbonate in, 198 carbonate in, 198 calcium oxide in, 198 calcium oxide in, 198 calcium oxide in, 198 calcium oxide in, 198 calculation of proteins, 153 carbonate in, 190 carbonate in, 198 calculation of proteins, 153 carbonate in, 198 calculation of proteins, 153 carbonate in, 198 calculation of proteins, 153 carbonate in, 198 calculation of proteins, 153 carbonate in, 198 calculation of proteins, 153 carbonate in, 198 calculation of proteins, 153 carbonate in, 198 calculation of proteins, 153 carbonate in, 198 cabonate in, 198 calculation of proteins, 153 carbonate in, 198 cabonate in,		ashing of, 134
chicory, 386 cinnamon, 426 cloves 416 cocoa, 403 cocoanut shells, 419 coffee. 386 corn, 309 starch, 281 date stones, 390 flour, 306, 322 ginger, 449 honey, 633 jams, 923 jellies, 923 lard, 557 mace, 466 milk, 124 mustard, 458 boric acid in, 182 calcium oxide in, 198 calculation of proteins, 153 caramel in, 176, 177 carbonate in, 180, 182 chocolate, 397 citric acid in, 127 carbonate in, 180, 182 calcium oxide in, 198 calculation of proteins, 153 caramel in, 176, 177 carbonate in, 180, 182 calcium oxide in, 198 calculation of proteins, 153 caramel in, 176, 177 carbonate in, 180, 182 chocolate, 397 citric acid in, 198 calculation of proteins, 153 caramel in, 176, 177 carbonate in, 180, 182 chocolate, 397 cohocolate, 397 cohoco	·	* * * * * * * * * * * * * * * * * * *
cinnamon, 426 cloves 416 cocoa, 403 cocoanut shells, 419 coffee. 386 corn, 309 starch, 281 date stones, 390 folur, 306, 322 ginger, 449 honey, 633 jams, 923 jellies, 923 lard, 557 mace, 466 milk, 124 mustard, 458 calcium oxide in, 198 calculation of proteins, 153 caramel in, 176, 177 carbonate in, 180, 182 chocolate, 397 citric acid in, 127 coloring matter in, 174–177 composition of, 124–126 constants, 169 ewe's, 127 fat of, 127, 134 fermentations of, 129 foods, prepared, 157 fore milk, 128 formaldehyde in, 178, 181 mosection, 159	cereal products, 305	hoiled milk, detection, 155
cloves 416 cocoa, 403 cocoanut shells, 419 coffee. 386 corn, 309 starch, 281 date stones, 390 fats, 510 flour, 306, 322 ginger, 449 honey, 633 jams, 923 jams, 923 lard, 557 mace, 466 milk, 124 mustard, 458 cocoanut shells, 419 carbonate in, 176, 177 carbonate in, 180, 182 chocolate, 397 citric acid in, 127 coloring matter in, 174–177 composition of, 124–126 constants, 169 ewe's, 127 fat of, 127, 134 fermentations of, 129 foods, prepared, 157 fore milk, 128 formaldehyde in, 178, 181 goat's, 127 human, 127 inspection, 159	chicory, 386	
cocoa, 403 cocoanut shells, 419 coffee. 386 corn, 309 starch, 281 date stones, 390 fats, 510 flour, 306, 322 ginger, 449 honey, 633 jams, 923 jams, 923 lard, 557 mace, 466 milk, 124 mustard, 458 caramel in, 176, 177 carbonate in, 180, 182 chocolate, 397 citric acid in, 127 coloring matter in, 174–177 composition of, 124–126 constants, 169 ewe's, 127 fat of, 127, 134 fermentations of, 129 foods, prepared, 157 fore milk, 128 formaldehyde in, 178, 181 goat's, 127 human, 127 inspection, 159	·	
cocoanut shells, 419 coffee. 386 corn, 309 starch, 281 date stones, 390 fats, 510 flour, 306, 322 ginger, 449 honey, 633 jams, 923 jams, 923 lard, 557 mace, 466 milk, 124 mustard, 458 chocolate, 397 citric acid in, 127 coloring matter in, 174–177 composition of, 124–126 constants, 169 ewe's, 127 fat of, 127, 134 fermentations of, 129 foods, prepared, 157 fore milk, 128 formaldehyde in, 178, 181 goat's, 127 human, 127 inspection, 159	cloves 416	calculation of proteins, 153
coffee. 386 corn, 309 starch, 281 date stones, 390 fats, 510 flour, 306, 322 ginger, 449 honey, 633 jams, 923 jellies, 923 lard, 557 mace, 466 milk, 124 mustard, 458 chocolate, 397 citric acid in, 127 coloring matter in, 174-177 composition of, 124-126 constants, 169 ewe's, 127 fat of, 127, 134 fermentations of, 129 foods, prepared, 157 fore milk, 128 formaldehyde in, 178, 181 goat's, 127 human, 127 inspection, 159	,	
corn, 309 starch, 281 date stones, 390 fats, 510 flour, 306, 322 ginger, 449 honey, 633 jams, 923 jellies, 923 lard, 557 mace, 466 milk, 124 mustard, 458 citric acid in, 127 coloring matter in, 174-177 composition of, 124-126 constants, 169 ewe's, 127 fat of, 127, 134 fermentations of, 129 foods, prepared, 157 fore milk, 128 formaldehyde in, 178, 181 goat's, 127 human, 127 inspection, 159		
starch, 281 date stones, 390 fats, 510 flour, 306, 322 ginger, 449 honey, 633 jams, 923 jellies, 923 lard, 557 mace, 466 milk, 124 mustard, 458 coloring matter in, 174-177 composition of, 124-126 constants, 169 ewe's, 127 fat of, 127, 134 fermentations of, 129 foods, prepared, 157 fore milk, 128 formaldehyde in, 178, 181 goat's, 127 human, 127 inspection, 159	=	
date stones, 390 fats, 510 fats, 510 flour, 306, 322 ginger, 449 honey, 633 jams, 923 jellies, 923 lard, 557 mace, 466 milk, 124 mustard, 458 composition of, 124–126 constants, 169 ewe's, 127 fat of, 127, 134 fermentations of, 129 foods, prepared, 157 fore milk, 128 formaldehyde in, 178, 181 goat's, 127 human, 127 inspection, 159		
fats, 510 flour, 306, 322 ginger, 449 honey, 633 jams, 923 jellies, 923 lard, 557 mace, 466 milk, 124 mustard, 458 constants, 169 ewe's, 127 fat of, 127, 134 fermentations of, 129 foods, prepared, 157 fore milk, 128 formaldehyde in, 178, 181 goat's, 127 human, 127 inspection, 159		
flour, 306, 322 ginger, 449 honey, 633 jams, 923 jellies, 923 lard, 557 mace, 466 milk, 124 mustard, 458 ewe's, 127 fat of, 127, 134 fermentations of, 129 foods, prepared, 157 fore milk, 128 formaldehyde in, 178, 181 goat's, 127 human, 127 inspection, 159	· · · · · · · · · · · · · · · · · · ·	
ginger, 449 honey, 633 jams, 923 jellies, 923 lard, 557 mace, 466 milk, 124 mustard, 458 fat of, 127, 134 fermentations of, 129 foods, prepared, 157 fore milk, 128 formaldeplde in, 178, 181 goat's, 127 human, 127 inspection, 159		
honey, 633 jams, 923 jellies, 923 lard, 557 mace, 466 milk, 124 mustard, 458 fermentations of, 129 foods, prepared, 157 fore milk, 128 formaldehyde in, 178, 181 goat's, 127 human, 127 inspection, 159		
jams, 923 foods, prepared, 157 jellies, 923 fore milk, 128 formaldehyde in, 178, 181 mace, 466 goat's, 127 human, 127 mustard, 458 inspection, 159		
jellies, 923 fore milk, 128 formaldehyde in, 178, 181 mace, 466 goat's, 127 human, 127 mustard, 458 foremalk, 128 formaldehyde in, 178, 181 goat's, 127 human, 127 inspection, 159		
lard, 557 formaldehyde in, 178, 181 mace, 466 goat's, 127 human, 127 mustard, 458 inspection, 159		
mace, 466 goat's, 127 milk, 124 human, 127 mustard, 458 inspection, 159		
milk, 124 human, 127 mustard, 458 inspection, 159		
mustard, 458 inspection, 159		• • •
nutmeg, 404 l known purity, 169	·	
	nutmeg, 404	known purity, 169

Wills manufacture	None and Waller was 4-11
Milk, mare's, 127	Munson and Walker sugar table, 599
methods of analysis, 130, 163, 163	Muscle albumin, 211
microscopical appearance, 124 modified, 155	fibers in meat, 211 sugar, 212, 238
nitrogen compounds in, 125, 145	Muscovado, 567, 568
powder, 157	Mustard, 453
preservatives in, 177	adulteration of, 459
proteins of, 125, 145, 153	ash of, 457
records of analysis of, 172	black, 455
references on, 208	cake, 455
ropy, 130	coloring matter in, 460
sampler, 131	composition of, 455, 456
serum, refraction of, 166, 167	Dakota, 460
specific gravity of, 166, 167	flour, 454
skimmed, 161	methods of analysis, 457
sour, analysis of, 186	microscopical structure of, 458
souring of, 129	oil, fixed, 454, 525
standards, 160	volatile, 453, 457
strippings, 128	pickles, gog
sucrate of lime in, 196	prepared, 460
detection, 197, 198	adulteration of, 460
sugar, 125, 577	composition of, 460, 461
determination of, 593, 594, 598	methods of analysis, 461
determination of, in milk, 147,	sinalbin in, 454
149, 151	mustard oil, 453
systematic examination of, 130, 168	sinapin sulphocyanate, 457
total solids in, 133, 134	standard, 459
calculation of, 151, 153, 154	starch in, 453
watering of, 161	turmeric in, 460
Milliau's cottonseed oil test, 518	volatile oil of, 453
Millon's reaction, 41, 92	wheat in, 459
reagent, 92	white, 453
Mill's bromine absorption method, 493	Mutton, composition of, 215
Mineral colors, 790	cuts of, 215
Mineral content of food, 47	tallow, 529
Mirbane, oil of, 875	Myosin, 42
Mitchell's fusel oil method, 748	insoluble, 44
Modified milk, 155	Natural min - 68 -
Mohler's test for benzoic acid, 829	Natural wine, 685
Moisture, determination of, 61 Molasses, 567	Neufchâtel cheese, 202
adulteration of, 621	Nickel salts, 899
ashing of, 614, 624	determination of, 903 Niebel's glycogen method, 236
clarifying, 614	Nitrates in food, 40, 46
composition of, 568	in watered milk, 168
glucose in, 621	Nitrobenzol, 875, 876
invert polarization at 87° C., 623	Nitrogen apparatus, 72, 73
methods of analysis, 613	compounds in milk, 145
standard for, 621	determination of, 69, 73
sucrose in, 614	free extract, 54
tin in, 625	Nitrogenous bodies
total solids in, 613	classification of, 40
vinegar, 763	separation of, in cheese, 205
Mollusks, 256	in meat, 228
Mucoid protein, 127	in milk, 125, 126, 145
Munson and Walker sugar method, 151, 598	Noodles, 347
=	ee • ▼

Notification, 10 Novau, 754 Nuclein, 43 Nucleoproteins, 43 Nutmeg, 462, 463 adulteration of, 464 composition of, 462, 463 extract, standards, 881 Macassar, 465 microscopical structure of, 464 oil of, 463 standard, 881 standard, 881 Nutrose, 158	Oils, edible, constants of, 508, 509 elaidin test, 499 fatty acids in, 484, 499 iodine absorption of, 487, 492 judgment as to purity of, 473 Maumené test, 494 melting point, 480 methods of analysis, 473 microscopical examination, 510 phytosterol in, 502, 503, 507 Polenske number of, 483 rancidity of, 473, 530 references on, 561 refractive index of, 477
Nuts, composition of, 275	Reichert-Meissl number, 481 saponification of, 472, 484, 486
Oats, 271 analysis of, 271, 272	sitosterol in, 522 specific gravity of, 474
ash of, 302	factors, 475
microscopic structure, 309	thermal tests, 493
starch in, 282	titer test, 500
Oil cakes, effects on butter of feeding, 531	unsaponifiable matter in, 501
lard of feeding, 560	Valenta test, 499
calorimeter, 495	viscosity, of 477
Oil, bitter almond, 873	Oils, essential, 880
cassia, 425	Oleomargarine, 541
cloves, 881	adulteration of, 543 coloring of, 542
cocoanut, 528 corn, 521	constants of, 544
cottonseed, 516	distinction from butter, 544, 546
ginger, 446	healthfulness of, 543
lard, 555	manufacture of, 541
lemon, 861, 871	microscopical examination, 552
terpeneless, 862	odor and taste, 545
lemongrass, 872	palm oil in, 542
mustard, fixed, 454, 525	Zega's test for, 553
volatile, 453, 457	Oleo oil, 541
nutmeg. 881	Olive, composition, 511
oleo, 541	Olive oil, 512
olive, 511	adulteration of, 512, 515
orange, 873	examination of, 515 refraction of, 514
peanut, 522	standard, 513
peppermint, 879 poppyseed, 526	Olives, pickled, 909
rape, 520	Olive stones, 436
rosin, 527	Orange colors, 785, 786, 806
sesame, 519	extract, 873
spearmint, 880	oil, 873
sunflower, 526	standards, 873
wintergreen, 878	terpeneless, 873
Oils, edible, 471. See also Fats	Orchil, 789
acetyl value, 497	O'Sullivan-Defren sugar method, 150
bromine absorption of, 492	Ovalbumin, 262
bromination test, 494	Oven, drying, 22
cholesterol in, 502, 503, 507	McGill's 586
composition of, 471, 472	Ovomucin, 262

0	D
Ovomucoid, 263	Pepper, olive stones in, 436
Oxygen absorbed, 415	piperin in, 429
equivalent, 415	determination of, 433
Oxyhæmoglobin, 43	red. See Cayenne and Paprika
Oysters, 257	shells, 435
	standard, 435
Palas rapeseed oil test, 521	varieties of, 429
Paprika, 439	white, 429
added oil in 445	Peppermint extract, 879
adulteration of, 444, 445	composition of, 880
composition of, 442	standards, 879
methods of analysis, 445	oil, 879
microscopical structure of, 441	Peptides, 45
Paraffin in confectionery, 647	Peptones, 44
in beeswax, 643	in cheese, 202
in fats, 510	in meat, 211, 231
in oleomargarine, 543	in milk, 146
Paranuclein, 206	Peter's test for benzoic acid, 829
Parenchyma, 87	Perry, 683
Pastes, adulteration of, 349	Persian berries, 788
artificial colors in, 349	Petroleum ether, 66
edible, 347	Phloroglucide, 286
Italian, 347	Phloroglucinol, 287
lecithin phosphoric acid in, 349	Phosphate baking powders, 333
methods of analysis, 349	Phosphoproteins, 43
noodles, 347	Phosphoric acid in baking chemicals, 346
Patrick's method for water in butter, 531	in beer, 725
test for thickeners in ice cream,	Phosphotungstic acid reaction, 45
200	Photomicrography, 93
Pea, composition, 272	camera for, 96
proteins of, 300	Phytolacca 788
starch of, 282	Phytosterol, 502
Peanut oil, 522	acetate test, 507
adulteration of, 523	crystallization of, 503
standards for, 522	determination of, 503
tests for, 523, 525	distinction from cholesterol, 503
Pear cider, 683	separation of, 503
essence, imitation, 884, 885	Piccalilli, 909
Pectose, 93, 276	Pickled meats, 218
Pekar's color test of flour, 317	Pickles, 909
Pentosans, 285, 296	adulteration of, 900
determination of, 285, 296	Pickling pump, 219
in cocoa products, 396	Pimiento, 439, 442
table for, 288	Pineapple essence, imitation, 884, 885
Pentose, 285, 296	Pioscope, 164
Pepper, 428	Piperin, 429
adulteration of, 435	determination of, 433
black, 429	Piutti and Bentivoglio's method for colors
buckwheat in, 437	in pastes, 351
composition of, 430, 432	Plant crystals, 90
dust, 436	Plasmon, 158
ether extract in, 410	
	Plastering, of wine, 692
long, 438	Platinum dishes, 61, 133, 134, 170
microscopical structure of, 433	counterweights for, 170
nitrogen in, 432	Poisoned foods, 74
in ether extract, 433	Poivrette, 436

•	
Polariscope, 578. See also Saccharimeter	Proteins, of milk, calculation of, 153
micro, 84	determination of, 145
Polariscope tube jacketted, 639	of peas, 300
short, for oils, 870	of potatoes, 301
Polarization at high temperature, 639	of rye, 277, 300
of essentional oils, 871	of wheat, 277, 298
honey, 639	secondary derivatives, 44
lemon extract, 864	simple, 41
molasses, 614	tests for, 41
orange extract, 873	Protein grains, 90
sugar, 578	Proteolytic fermentation, 158, 202
vinegar, 769	Proteoses, 44, 297
Polarization of wine, 694, 703	Proto-albumose, 44, 45
Polenske number, 483	Proximate analysis, extent of, 53
Poppyseed, 526	expression of results of,
oil, 526	53
Pork, composition of, 216	Prussian blue, 790, 811
cuts of, 216	Ptomaines, 218
Porter, 709, 712. See also Beer.	Publication of adulterated foods, 10
Port wine, 689	Pulfrich refractometer, 100
Potash determination, 304, 345	Pycnometer, 57
Potassium myronate, 453, 457	Pyroligneous acid, 764
Potatoes, composition of, 273	
proteins of, 301	Quassiin, 727
starch of, 282	Quercitannic acid, 415
Poultry, composition of, 216	Quevenne's lactometer, 132
Preparation of sample, 55	Quince essence, imitation, 884, 885
Preservatives, 815	Quotient of purity of sugar, 586
commercial food, 817	
in butter, 538	Raffinose, 279, 577
in canned goods, 903	determination of, 620
in fish, 257	Rancidity, 473, 530
in meats, 220, 232	Rape oil, 520
in milk, 177, 183	test for, 521
in table sauces, 908	seed, 520
of eggs, 266, 268	Raphides, 90
references on, 838	Reagents, 35, 90
regulation of, 816	references on, 38
Pressure pump, 20	table of, 26-34
Process butter, 540	Red colors, 783, 784, 788, 792, 804
Prolamins, 42	Red ochre in sausages, 238
Proof spirit, 677	Red pepper. See Cayenne and Paprika
Prosecution, 10	Red wines, 684, 689
	1 = 1
Protamins, 43	Red wood, 444
Proteans, 44	References on beer 756
Proteins, 40	butter, 562
coagulated, 44	canned goods, 927
conjugated, 43	cereals, 361
derived, 44	сосоа, 406
factor for, 40	coffee, 406
of barley, 277, 300	colors, 813
of beer, 725	dietetics, 49
of cereals, 296	distilled liquors, 758
of condensed milk, 190	eggs, 270
	flavoring extracts, 886
of eggs, 262	flesh foods, 258
of milk, 125	nesii ioods, 250

References on food economy, 49	Rose, extract, standards, 882
inspection, 11	rose oil in, 883
fruit products, 927	Rosin oil, 527
fruits, 361	Rota's color scheme, 797
general analytical methods, 79	Rubner's fuel value factors, 48
laboratory equipment, 38	Rum, 742
leavening materials, 364	composition of, 742
liquors, 756	essence, 743
microscope, 98	methods of analysis, 745
milk, 208	new, 743
oils, 561	standards, 742
preservatives, 838	Rye, composition of, 271
reagents, 38	microscopical structure of, 308
refractometer, 122	proteins of, 300
spices, 468	starch, 281
sugars, 650	Saccharimeter, 578
tea, 406	double wedge, 581
. ' '	forms of, 583
vinegar, 778	
wine, 757	normal weights for, 583
Refractometer, 100	scales compared, 583
Abbé, 100	single wedge, 579
Amagat and Jean, 100	Soleil-Ventzke, 578
butyro, 100, 101	triple shadow, 581
heater for, 102	Saccharimetry, 578
immersion, 111	Saccharin, 842
in oil analysis, 477	detection of, 843
Pulfrich, 100	determination of, 844
sliding scale for, 107	Saccharine products, 565
tables for, 104, 105, 113,	Saccharoses, 565
116, 120, 121	Safflower, 789
Wollny, 100, 139	Saffron, 789
Reichert-Meissl method, 481	Sago, 283
Reichert number of butter, 549	Saleratus, 332
Reinsch's test for arsenic, 728	Salicylic acid, 825
Relishes, 906	detection of, 825
Renard's test for peanut oil, 523	determination of, 826
for rosin oil, 527	in meat, 233
Renovated butter, 540	in milk, 180
distinction from butter	Salmin, 43
and oleomargarine, 546	Salted meats, 218
Resins, 89	Sample, preparation, 55
Respiration calorimeter, 2	Sanatogen, 158
Rice, composition of, 272	Sanose, 158
microscopical structure of, 310	Saponification, 472, 484, 486
starch, 282	Sarcolemma, 211
Riche and Bardy methyl alcohol method,	Sausages, 223
751	ash of, 225
Richmond's cane sugar method, 185	color of, 224
sliding milk scale, 153	composition of, 223
Ritsert's tests for acetanilide, 859	fat in, 226
Ritthausen's method for milk proteins, 145	glycogen in, 234
Roeser's mustard oil method, 457	horseflesh in, 234
Ropy milk, 130	methods of analysis, 225
Roquefort cheese, 202	starch in, 223
Rose, attar of, 882	water in, 225
extract, 882	Sauterne wine, 685, 688
January 00-	,,,

•	
Savory extract, standards, 881 oil, standards, 881	Specific gravity of milk serum, 166 temperature cor-
Sawdust, 450	rection for 133
Schied m schnapps, 744	of oils, 474
Schenk beer, 708	of vinegar, 764
Schlegel's method for colors in pastes, 350	
Schreiner's colorimeter, 77	Spent tea leaves, 375
Schultze's reagent, 93	Spices, 408
	- · · · · · · · · · · · · · · · · · · ·
Scienchyma, 87	adulterants of, 413
Scovell sampling tube, 131	alcohol extract of, 410
Sealed samples, 6, 159	ash of, 409
Semolina, 347	crude fiber of, 411
Separatory funnel support, 68	ether extract of, 410
Sericin, 42	lime in, 410
Sesame oil, 518	methods of analysis, 408
adulteration of, 519	microscopical examination of, 412
tests for, 519	nitrogen in, 410
seeds, 518	references on, 468
Sherry wine, 687	starch in, 411
Short's method for fat in cheese, 205	volatile oil of, 411
Shredded wheat, 352	Spiral ducts, 89
Sieve tubes, 89	
	Spirits, cologne, 731
Silent spirit, 731	distilled, 730
Sinabaldi's asaprol method, 838	neutral 731
Sinalbin, 454	silent, 731
mustard oil, 454	standards, 730
Sinigrin, 453	velvet, 731
Sinks, 17	Spirit vinegar, 760, 763
Sitosterol, 522	Spoon test for butter, 549
Smoked meats, 218	Sprengel tube, 60
"Soaked," goods, 905	Stahlschmidt's caffeine method, 374
Soda, determination of, 304, 345	Standards for allspice, 424
Sodium benzoate, 827	anise extract, 880
bicarbonate, 332	oil, 880
bisulphite, 833	beer, 711
carbonate, in milk, 180, 182	brandy, 740
hydroxide, tenth-normal solution, 35	butter, 535
salicylate, 825	cassia, 428
Soja bean meal, 357	extract, 880
Soleil-Ventzke saccharimeter, 578	oil, 880
Sorghum, 573	cayenne, 443
Sostegni and Carpentieri's test, 794	celery seed extract, 880
Souring of milk, 129	oil, 880
• •	•
Sour milk, 139	cheese, 203
Soxhlet extractor, 63	cinnamon, 428
Soxhlet's milk sugar method, 150, 152	extract, 881
Spaghetti, 347	oil, 881
Sparkling wine, 685, 691	clove extract, 881
Spearmint, extract, 880	oil, 881
standards, 880	cloves, 418
oil, 880.	cocoa, 402
Specific gravity bottle, 57	cream, 195
of beeswax, 643	foods, 4
of liquids, 55	ginger, 450
of liquors, 657	extract, 881
of milk, 131	oil, 88x
	•

_	•
Standards for ice cream, 199	Starch, pea, 282
lard, 556	potato, 282
lemon extract, 861	rice, 282
oil, 861	rye, 281
mace, 460	sago, 283
maple products, 572	syrup, 575
meats, 218	tapioca, 282
meat extracts, 241	under polarized light, 283
milk, 160, 162	wheat, 281
molasses, 621	Stearin, beef, 541
mustard, 459	cottonseed, 517
nutmeg, 464	lard, 555
extract, 881	Sterilized butter, 540
oil, 881	Still. alcohol, 659
olive oil, 513	fractionating, 67
pepper, 435	nitrogen, 73
renovated butter, 541	water, 22
rum, 742	wine, 685
savory extract, 881	Stilton cheese, 202
oil, 881	Stokes' milk centrifuge, 136
staranise extract, 881	Stone's method of carbohydrate separation,
oil, 881	295
starch sugar, 574	Storch's method for boiled milk, 155
sugars, 566, 574, 772	mucoid protein, 127
sweet basil extract, 881	Stout, 709, 712. See also Beer
oil, 88r	Strippings, 128
majoram extract, 881	Stutzer's gelatin method, 231
oil, 881	Suberin, 89
thyme extract, 881	Sucrate of lime, 196
oil, 881	Sucrose. See Cane sugar
vanilla extract, 853	Suction pump, 19
vinegar, 770	Suet, 529
wine, 689	Sugar, 561
whiskey, 733	beet, 569
Standard solutions, equivalents of, 36 refractometric readings	brown, composition of ash, 567
of, 120	cane, 566, 567
Staranise extract, standards, 881	classification of, 565 composition of, 568
oil standards, 881	grape. See Dextrose
Starch, 47, 89, 279	in fruits, 566
arrowroot, 282	maple. See Maple syrup
barley, 281	methods of analysis, 585
bean, 282	muscovado, 567
buckwheat, 281	organic non-sugars in, 586
classification of, 280	quotient of purity, 586
corn, 281	raw, 568, 569
detection of, 279	references on, 650
determination of, 283	refining, 570
by acid conversion, 283	standards, 566, 572, 574
by diastase method, 283	ultramarine in, 570, 590
in baking powder, 343	Sulphur, determination of, 305
in cereals, 283, 296	Sulphuric acid in baking chemicals, 346
in milk, 185	in vinegar, 767
in sausages, 233	Sulphuring, 833
in spices, 411	Sulphurous acid, 833
oat, 282	detection of, 834
,	,

932	
Sulphurous acid, determination of, 834	Tin, action of fruits and vegetables on, 892,
in meat, 220, 232	893, 895
Sunflower oil, 526	determination of, 900, 902
seeds, 527	salts in molasses, 625
Sweet basil extract, standards, 881	Tintometer, Lovibond, 78
oil, standards, 881	Titer test, 500
Sweeteners, artificial, 842	Tocher's sesame oil test, 519
Sweet majoram extract, standards, 881	Tomato ketchup, 906
oil, standards, 881	coloring in, 907, 908
Sweet wine, 685, 690	preservatives in, 908
Syrup, analysis of, 613	Tonka bean, 852
ashing of, 614	tincture, 853
maple. See Maple syrup	Trillat methyl alcohol test, 750
mixing, 576	Turmeric, 450
starch, 576	as an adulterant, 452
total solids in, 613	microscopical structure of, 45r
Sy's lead method, 630	tests for, 453, 789
	Ultramarine blue, 791, 810
Table sauces, 905	in sugar, 570, 590
preservatives in, 908	Uno beer, 714
Tallow, 529	Unsaponifiable matter, 501
Tannin in cloves, 415	c isaponimusic muteer, 301
in tea, 370	Vacuoles in yeast cells, 330
in wine, 704	Vanilla bean, 849, 850
Tapioca, 282	exhausted, 851
Tartaric acid in baking powder, 339, 340	Vanilla extract, 849
in fruit products, 920	adulteration of, 853
Tartrate baking powders, 332	alcohol in, 860
Tea, 365	· alkali in, 852
adulteration of, 374	artificial, 854
ash of, 368, 369	caramel in, 860
astringents in, 377	color quotient of, 861
caffeine in, 372, 373	composition of, 851
composition of, 366, 367	coumarin in, 854
exhausted leaves in, 375	determination of, 858;
extract of, 370	glucose in, 860
facing of, 374	glycerin in, 860
foreign leaves in, 376	methods of analysis, 855
leaf, characteristics of, 376	prune juice in, 854
methods of analysis, 368	resins in, 855
microscopical examination of, 378	standards, 853
references on, 406	tannin in, 856
spent leaves in, 375	tonka in, 854
stems in, 376	vanillin in, 856, 858
tablets, 377	Vanillin, 851
tannin in, 370	determination, 856, 858
theine in, 372, 373	microscopical structure, 860
Tecnique, 82	Van Slyke's protein formula, 153
Teller's method of separating wheat pro-	method of nitrogen separations
teins, 298	in cheese, 205, 206
Theine, 372	in milk 146
Theobromine, 396, 400	Vaporimeter, 675
Thompson's boric acid method, 823	Veal, composition of, 214
Thyme extract, standards, 881	cuts of, 214
oil, standards, 881	Vegetable colors, 787

INDEX.

1	Viengar, standards, 770
Vegetable colors, in sausages, 239	sugar, 771
Vegetables, 273	sugars in, 769, 774
ash, of 302	sulphuric acid in, 767
composition of, 273	tartrate in, 769
methods of proximate analy-	tests on, 775
sis of, 276	varieties of, 759
references on, 361	volatile acids of, 76
Ventilation, 15	wine, 761, 771
Vermicelli, 347	wood, 764, 777
Vessels, 89	zinc in, 777
Villivecchia and Fabris' sesame oil test, 520	Vinous fermentation, 654
Vinegar, 759	Viscogen, 196
acidity of, 765	Viscosity of cream, 196
acids of, 766	of oils, 477
adulterated, 776	Vitellin, 43
adulteration of, 770	
alcohol in, 766	Waage's Bombay mace test, 408
apple, 771	Walnut ketchup, 907
arsenic in, 778	Water-bath, 21
artificial, 772	Water glass, 266
ash of, 761, 764, 773 solubility and alkalinity of,	Waterhouse butter test, 550
	Weiss beer, 709
764	Werner-Schmidt method for fat in cheese, 205
beer, 762	in milk, 139
caramel in, 777	Wastabal balance #6
cider, 760. 771 artificial, 772	Westphal balance, 56
7.7%	West's benzoic acid method, 832
composition of, 760	Wheat, 271, 272
copper in, 778	ash of, 302
distilled, 763, 771 extract of, 764	composition of, 271, 272 microscopic structure of, 306
furfural in, 777	proteins of, 277, 298
glucose, 763, 771	shredded, 352
grain, 771	starch, 281
Hortvet number of, 768	Whiskey, 731. See also Distilled liquors
hydrochloric acid in, 767	adulteration of, 738
lead in, 777	aging of, 732
acetate, test for, 768, 777	American, 735
number of, 768	Bourbon, 732, 734, 736, 737
levulose in, 770	British, 735
malic acid in, 767	composition of, 734
malt, 762, 771	imitation, 738
manufacture of, 760	Irish, 732, 734, 735
metallic impurities in, 777	manufacture of, 731
methods of analysis, 764	methods of analysis, 745
mineral acids in, 766, 767	rye, 732, 734, 737
molasses, 763	Scotch, 732, 734, 735
nitrogen in, 765	standards, 733, 734
phosphoric acid in, 764	Wijs's iodine absorption method, 492
polarization of, 769, 774	Wild's saccharimeter, 583
reducing sugars in, 770	Wiley's bromine pipette, 495
references on, 778	Wiley and Ewell's double dilution sugar
residue of, 772	method, 149
specific gravity of, 764	Wine, 684
spices in, 777	acidity of, 696
spirit, 771	added alcohol in, 695

Wine, adulteration of, 601 ameliorated, 601 Burgundy, artificial, 692 California, 688 cane sugar in, 603 Cazeneuve's color method, 705 chaptalizing, 693 claret, 687 artificial, 692 classification of, 685 coloring matter in, 704, 705 composition of, 686 corrected, 691 cream of tartar in, 702 "dry," 690 Dupré's color method, 705 extract in, 696, 697 fortified, 685, 690 fruit other than grape, 695 glycerin in, 703 hocks, 689 Madeira, 685, 686 Malaga, artificial, 692 malic acid in, 702 manufacture of, 684 methods of analysis, 696 modified, 691 natural, 685 non-volatile acids in, 701 plastering, 692 polarization of, 703 port, 689 potassium sulphate in, 704 raisin, 691 red, 684, 689 reducing sugar in, 703 references on, 757 sherry, 687 artificial, 692 sparkling, 685, 691 standards, 689 still, 685 sweet, 690 tannin in, 704 tartaric acid in, 701 varieties of, 687

Wine, vinegar, 761 volatile acids in, 606 watering of, 604 white, 684, 689 yeast of, 684 Wintergreen extract, 878 adulteration of, 878 wintergreen oil in, 878 oil of, 878 Winton lead number, 628, 768 Wollny milk fat refractometer, 100, 139 tables for using, 141 table for converting Wollny degrees into nD, 143 Woodman and Newhall's color quotient, 861 and Taylor's caffetannic acid method, 383 Wood vinegar, 764, 777 Wool, double dyeing method with, 793 dyeing of, 793 for color tests, 793 vegetable colors on, 795 Xanthin, 46, 211 Xantho-proteic reaction, 41 Xylan, 285, 288, 296 Xylose, 285, 288, 296 Yeast, 327 adulteration of, 331 composition of, 329 compressed, 328 dry, 328 in cider, 678 in wine, 684 microscopical examination of, 320 starch in, 331 testing, 330 vacuoles in, 330 Yeast extracts, 246 Yellow colors, 783, 785, 788, 792, 806 Zega's test for oleomargarine, 553 Zein, 42, 300 Zinc salts, 897

determination of, 900

Fig. 121.—Barley, ×110.

Transverse section, showing in order, pericarp, seed coats, alcurone layer, and starch cells.

Fig. 122.—Barley, ×55.
Surface view of epidermis with hairs.

Fig. 123.—Barley, ×125.
Surface view of upper chaff layer.

Fig. 124.—Barley Starch, ×220.

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Fig. 125.—Buckwheat, ×110.

Transverse section through part of pericarp, seed coat, and part of endosperm.

Fig. 126.—Buckwheat, ×110. Surface view of scutellum.

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Fig. 129.—Buckwheat Starch, X110. Starch grains in masses. Fig. 130.—Corn, ×110.

Transverse section through pericarp, seed coat, proteid layer, and part of endosperm, showing starch cells.

Fig. 131.—Corn, ×110.
Surface view showing two layers of the mesocarp.

Fig. 132.—Corn, X 110. Surface section. Proteid layer.

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Fro. 133.—Com Starch, ×220.

Fig. 134.—Corn Starch, ×220. . With polarized light.

Fig. 135.—Oat, X110.
Transverse section through chaff.

Fig. 136.—Oat, ×110.

Surface section. Proteid layer with fragments of epidermis and hairs.

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Fig. 137.—Oat, ×110. Surface view of upper chaff layer. Fig. 138.—Oat, ×55. Surface view of epidermis and hairs.

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Fig. 141.—Rice, ×110. Surface section through starch cells. FIG. 142.—Rice, ×110. Surface view of upper chaff layer



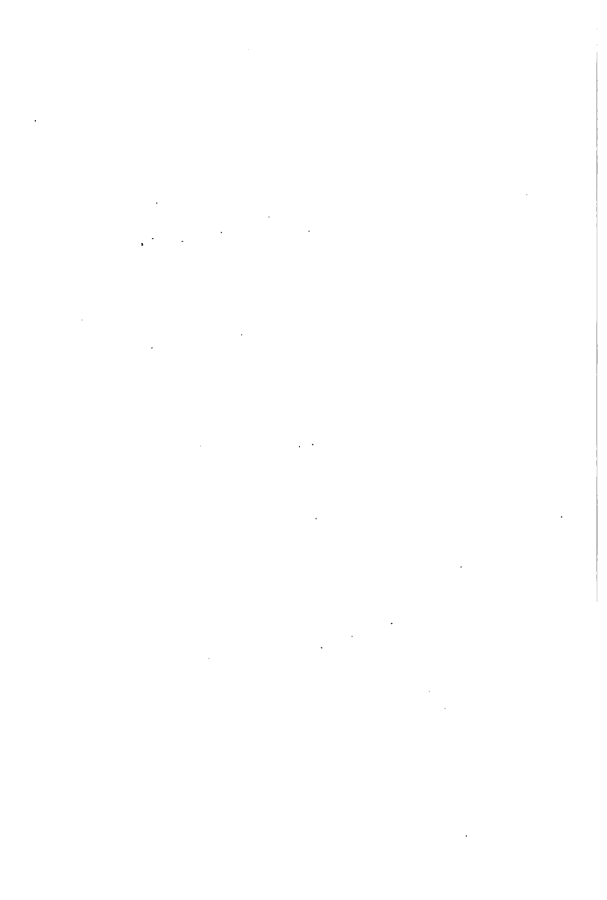
Fig. 145 —Rye, ×110.

Transverse section through pericarp, seed coat, aleurone layer, and starch cells of endosperm.

Fig. 146.—Rye, ×110 Surface view of epidermis and underlying layers.

Fig. 147.—Rye, ×110.
Surface view of epidermis and of seed cons.

Fig. 148.—Rye Starch, ×220.



#### PLATE VIII.

#### CEREALS.

Fig. 149.—Wheat, ×110.

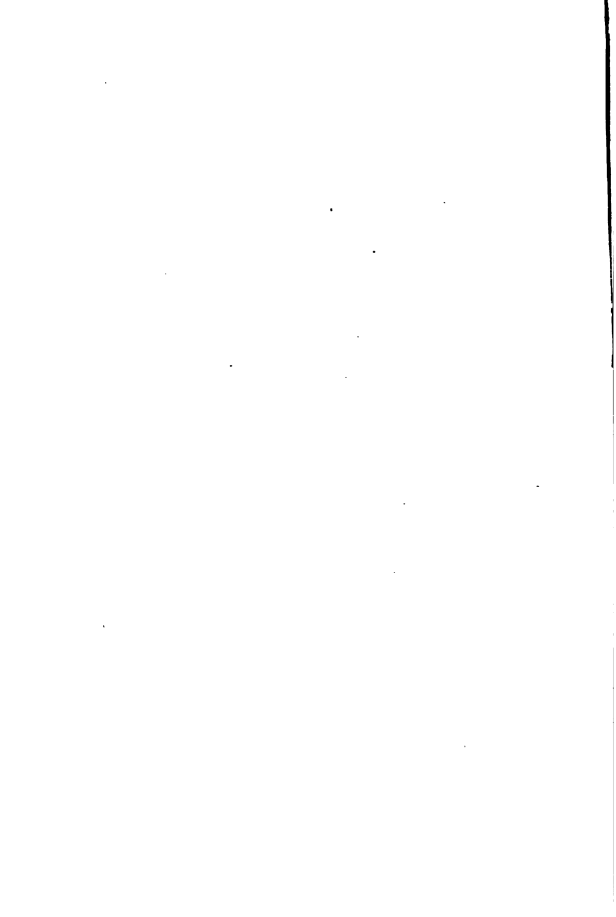
Transverse section through pericarp, seed coat, proteid layer, and starch cells of endosperm.

Fig. 150.—Wheat, × 110.

Surface view of outer and inner epidermis Also showing proteid layer.

Fig. 151.—Wheat, ×110. Surface view of epidermis, with hairs.

FIG. 152.—Wheat Starch, ×220.



#### PLATE IX.

## LEGUMES.

Fig. 153.—Bean, × 110. Transverse section through starch cells. Fig. 154.—Bean Starch, ×220.

Fig. 155.—Bean, X110.

Transverse section through hull, showing palisade cells of epidermis, and underlying hypoderma.

Fig. 156.-Lentil, X 110.

Transverse section through hull and part of endosperm, showing some of the starch cells.

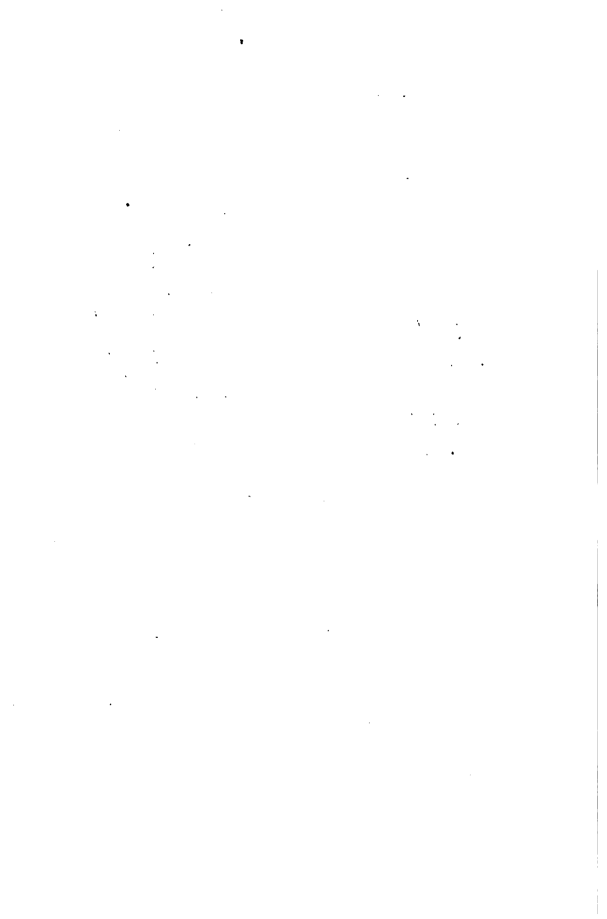


PLATE X.

LEGUMES.

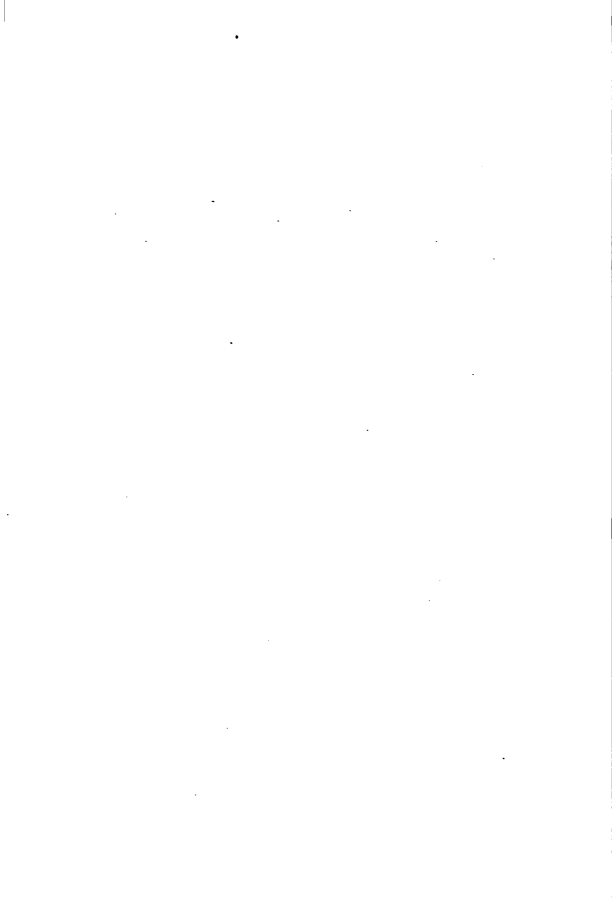
Fig. 157 —Lentil, ×110. Surface view of epidermis.

Fig. 158.—Pea, ×110.

Transverse section through hull and seed coat, showing outer palisade cells and underlying hypoderma.

Fig. 159.—Pea. ×110.
Surface section through base of palisade layer.

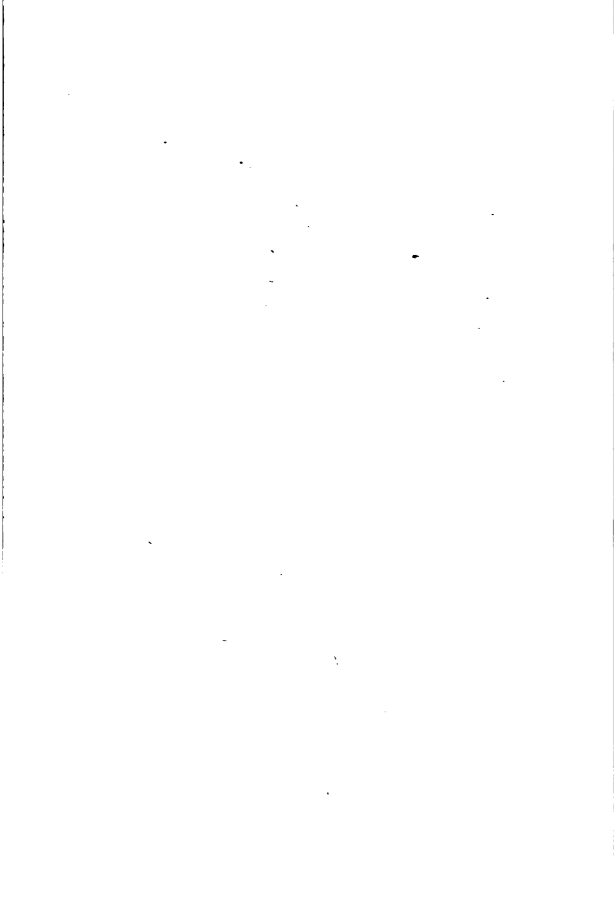
Fig. 160.—Pea, X110.
Powdered pea hulls.



## LEGUMES.

Fig. 161.—Pea, ×110. Surface view of palisade cells.

Fig. 162.—Pea, ×110. Transverse section through starch cells.



### PLATE XII.

# MISCELLANEOUS STARCHES.

Fig. 165.—Potato Starch, ×220.

Fig. 166.—Potato Starch, ×220. With polarized light.

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#### PLATE XIII.

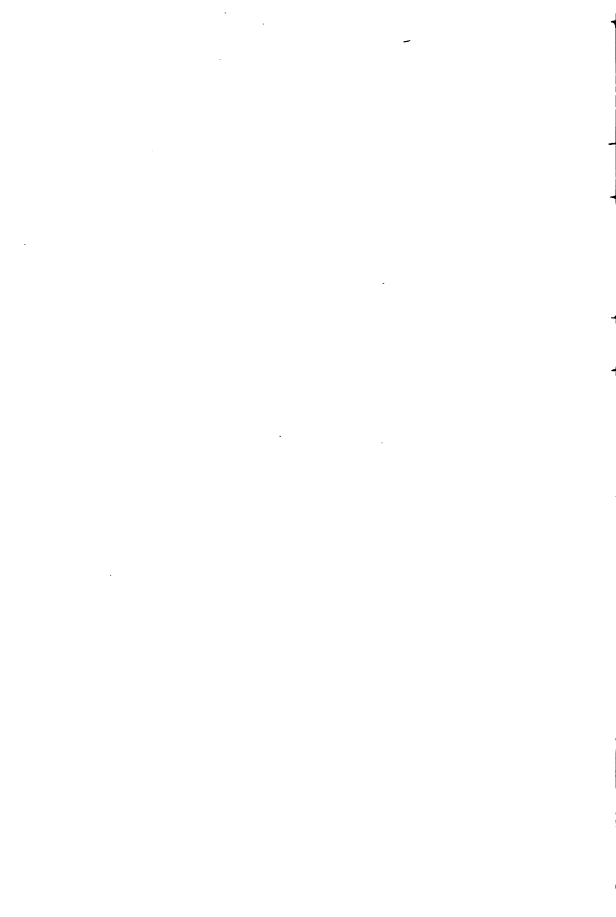
# TURMERIC. SAGO.

Fig. 169.—Turmeric, × 70.

Transverse section through rhizome.

Fig. 170.—Turmeric, ×110.

Longitudinal section. Note spiral ducts through the center.



COFFEE.

F10. 173.—Raw Coffee, ×110.

Transverse section of outer portion of endosperm.

Fig. 174.—Roasted Coffee, ×130.

Transverse section through parenchyma of endosperm.

Fig. 175.—Coffee, X110. Surface view of seed coat. Fig. 176.—Coffee, XIIc

Roasted, ground coffee, showing fragments of endosperm parenchyma and of seed coat.

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PLATE XV.

COFFEE. CHICORY.

Fro. 177.—Adulterated Coffee, ×130.

Dark masses of roasted pea starch are shown, with transparent fragments of the palisade cells of the pea-hull.

Fig. 178.—Adulterated Coffee, × 130.

The vascular ducts of chicory show most conspicuously in this field.

Fig. 179.—Chicory, ×25.
Transverse section through the root.

Fig. 180. -Chicory, X 110. Transverse section.

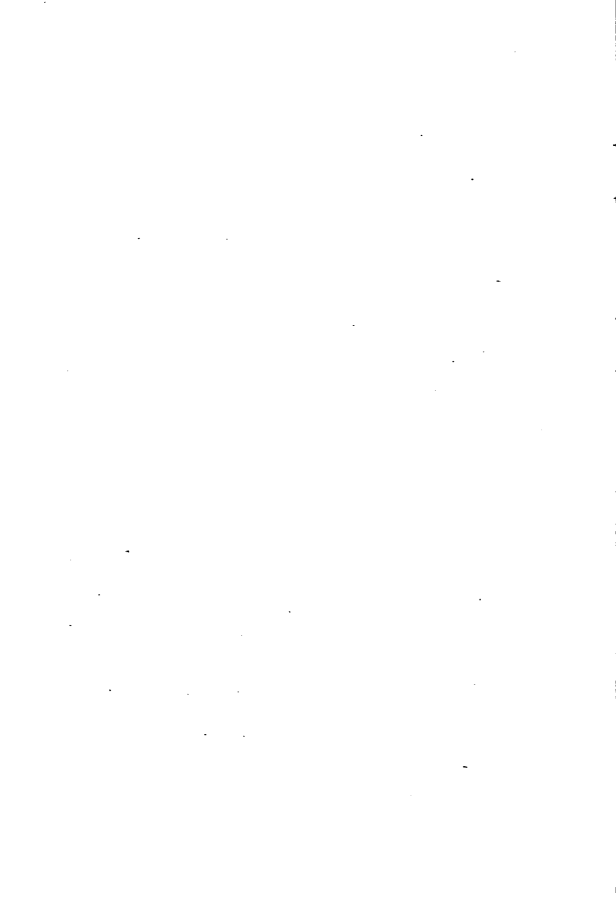


PLATE XVI.

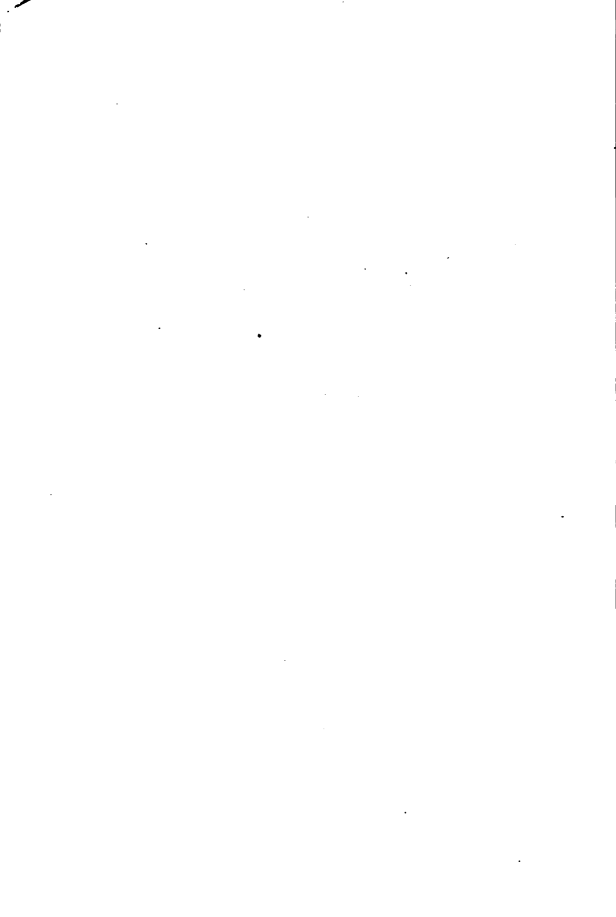
CHICORY. COCOA.

Fig. 181.—Chicory, X110.

Fig. 182.—Chicory, X110 Tangential section, showing reticulated ducts and Radial section, showing bark parenchyma and wood parenchyma. Radial section, showing bark parenchyma and milk ducts.

Fig. 183.—Chicory, ×110. Roasted and ground, showing fragments of ducts and other tissues.

Fig. 184.—Cocoa, ×110. Transverse section through periphery of seed, seed coats, and cotyledon.



COCOA.

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Fig. 185.—Powdered Cocoa, X110.

Fig. 186.—Adulterated Cocoa, ×110 Showing admixture of arrowroot with the cocoa powder.

Fig. 187.—Cocoa Shell, ×110.

Transverse section through epidermis, pulp, and mucilaginous layers of the pericarp and seed coat.

Fig. 188.—Cocoa Shell, × 110.
Longitudinal section through shell.



TEA. SPICES.

Fig. 189.—Tea, ×55.

Transverse section through midrib of leaf. Note the palisade layer below the upper epidermis, the inner wood vessels above the center, and the parenchyma of the pulp. Fig. 190.—Tea, × 110.
Surface view of lower epidermis, with stomata and one of the hairs.

Ftg. 191.—Allspice, ×9.

Transverse section through the entire berry, showing the two cells, with kidney shaped seed in each. Fig. 192.—Allspice. × 70.

Transverse section through pericarp, showing oil spaces and stone cells.



Fig. 193.—Allspice Seed ×110.

Transverse section through seed shell and part of embryo, showing starch cells.

Fig. 194.—Allspice Seed, X110.

Transverse section through the resinous portion of the seed coat, showing port wine colored lumps of gum or resin.

Fig. 195.—Powdered Allspice, XIIO. Showing stone cells, resinous lumps, and starch. Fig. 196.—Adulterated Allspice, ×110.

Showing a large fragment of the seed skin of cayenne at the left.

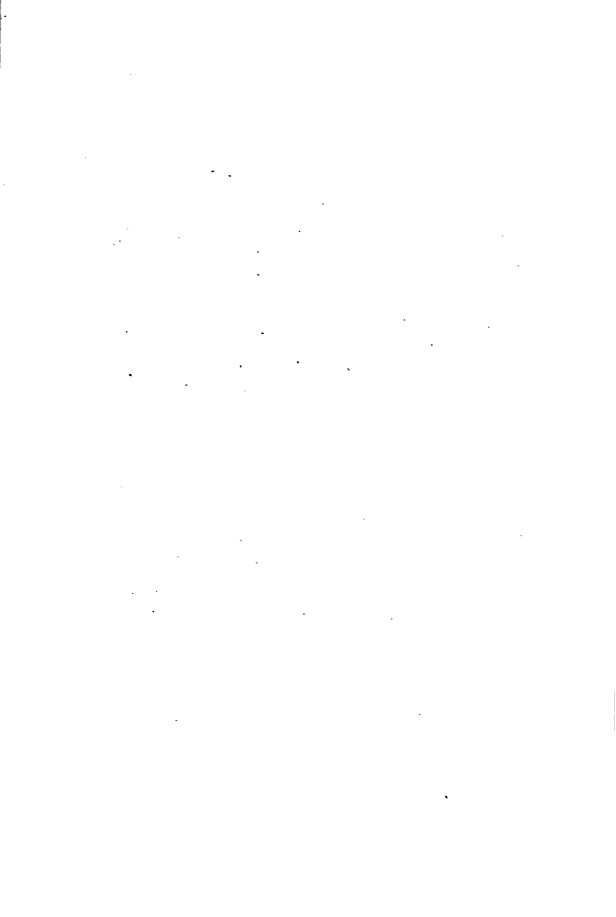


Fig. 197.—Cassia Bark, ×45.
Transverse section through the bark.

Fig. 198.—Cassia Bark, ×45. Longitudinal section.

Fig. 199.—Cassia Bark, ×110.

Transverse section, showing cork cells, parenchyma, and stone cells.

Fig. 200.—Cassia Bark, ×110.

Longitudinal section, showing bunches of bast fibers at the left, starch cells in the center, and stone cells at the right.



Fig. 201.—Ceylon Cinnamon Bark, ×110. Transverse section, showing many bast fibers and starch cells. Fig. 202.—Ceylon Cinnamon Bark, X110.

Longitudinal section, showing bast fibers, stone cells, and parenchyma.

Fig. 203.—Powdered Cassia, X110. Showing stone cells, starch, and corky tissue. Fig. 204.—Powdered Cassia, X110. Showing bast fibers and starch.



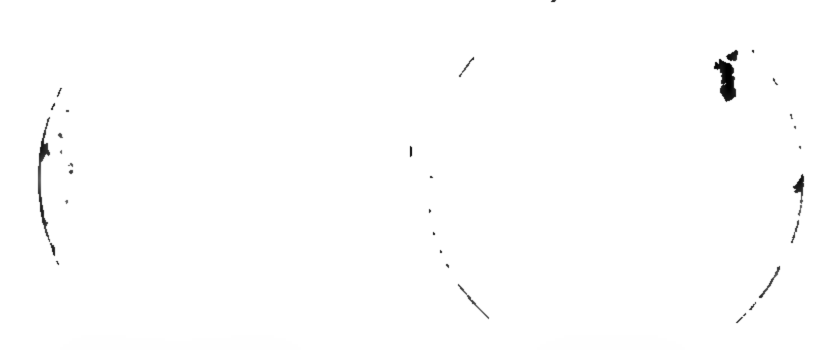


FIG. 205.—Powdered Cassia, × 110. "nowing large bast fiber and starch grains.

Fig. 206.—Adulterated Cassia, × 110.
A mass of foreign bark.

Fig. 207.—Cayenne, ×110. Transverse section through pericarp. Fig. 208.—Cayenne, × 110.

Transverse section through seed coat and part of endosperm. Collapsed parenchyma cells separate endosperm from long epidermal cells.



## PLATE XXIII.

SPICES.

Fig. 209.—Cayenne, X110. Surface view of fruit epidermis.

Ftg. 210.—Cayenne, X110. Surface view of two layers of seed coat.

Fig. 211.—Powdered Cayenne, ×110.
A large mass of fruit epidermis.

Fig. 212.—Powdered Cayenne, ×110. Showing chiefly two of the seed coat layers.



Fig. 213.—Adulterated Cayenne, ×130. Corn and wheat starch and cocoanut shells appear chiefly. A bit of cayenne is shown at the right.

Fig. 214.—Adulterated Cayenne, ×214. The central mass is ground red wood, surrounded by corn starch grains.



Fig. 217.—Clove, ×28.

Longitudinal section through entire clove.

Fig. 218.—Clove, ×70. Central longitudinal section, showing duct bundles.

Ft6 219.—Clove, ×110. Surface view of epidermis.

Fig. 220.—Powdered Cloves, X130.

Dense, spongy tissue, with small oil drops.

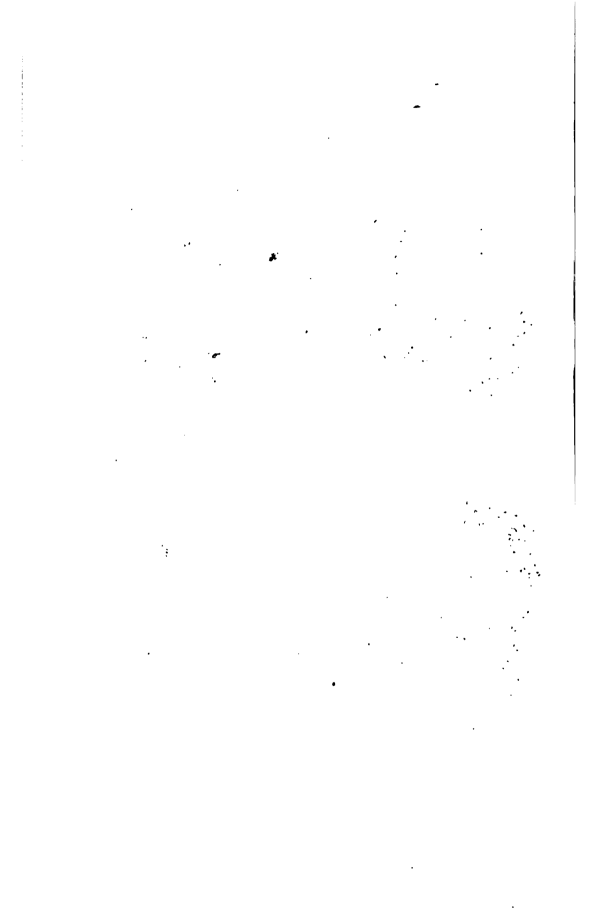


Fig. 221.—Clove Stem, ×70.

Transverse section through outer part of stem, showing bast fibers at the left, parenchyma in the center, and stone cells near the epidermis.

Fig. 222.—Clove Stem, ×25. Central longitudinal section through entire stem, showing bast fibers in the center, and stone cells at the right.

Fig. 223.—Clove Stem, ×70. Longitudinal section, showing the stone cells.

FIG 224.—Powdered Clove Stems, XIIO. Showing fragments of tissues, stone cells, and bast fibers.

Fig. 225.—Powdered Clove Stems, ×110. Showing bundle of bast fibers. Ftc. 226.—Adulterated Cloves, ×130. Showing chiefly stone cells of cocoanut shells.

Fig. 227.—Adulterated Cloves, ×130. With large admixture of cocoanut shells.

FIG. 228.—Ginger, XIIO.

Transverse section, showing starch cells with contents.



Fig. 229.—Ginger, ×110.

Fig. 230.—Ginger, ×110. Transverse section, showing parenchyma, starch grains, and duct vessels.

Longitudinal section, showing spiral ducts and pigment cells.



Fig. 231.—Ginger Starch, ×220.

Fig. 232.—Adulterated Ginger, X 130. A mass of wheat bran tissue is most conspicuous.



Fig. 233.—Adulterated Ginger, ×130. The central dark mass is a yellow fragment of Containing a large admixture of corn and wheat turmeric.

Fig. 234.—Adulterated Ginger, ×130. starches.

Fig. 235.—Penang Mace, X110. Transverse section through epidermis and oil cells, showing also parenchyma with contents of amylodextrin.

Fig. 236.—Bombay or Wild Mace, X110. Transverse section through outer layers, showing yellow and red resinous lumps.

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Fig. 237.—Nutmeg, ×110.

Transverse section through the exterior and interior teguments of the seed and part of the endosperm, showing starch cells.

Fig. 238.—Nutmeg, ×25. Transverse section near exterior of seed.

F16 239.—Nutmeg, × 110.

Surface view of seed coat, showing also portions of underlying tissues.

Fig. 240.—Powdered Nutmeg, X110.

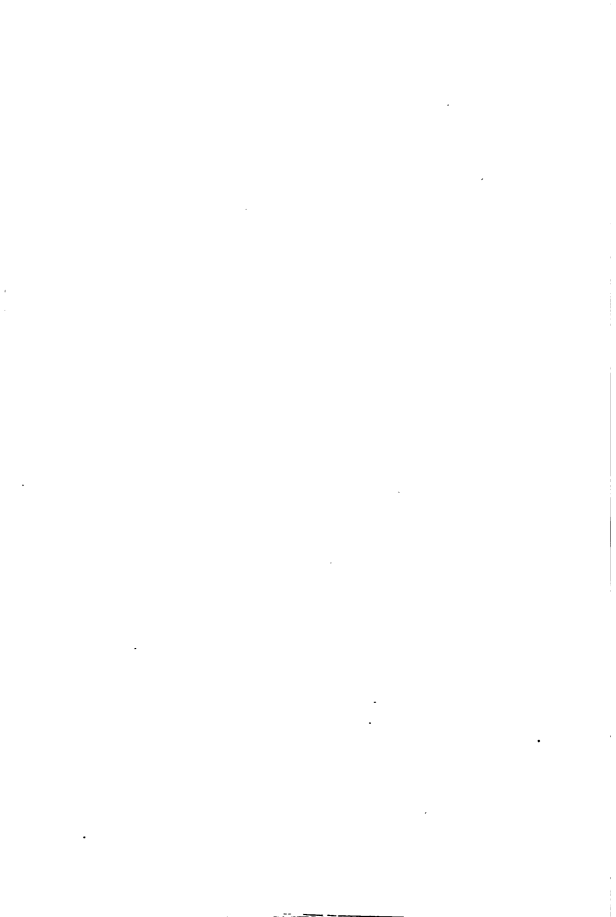


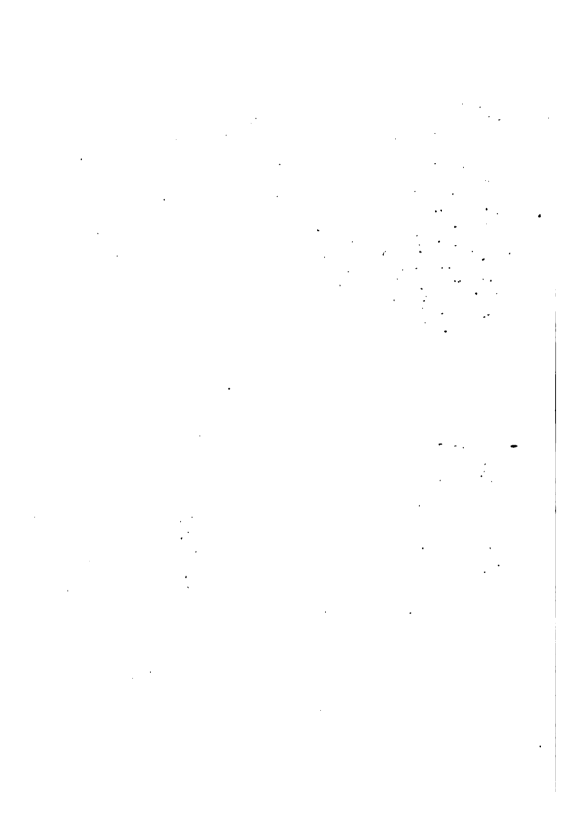
Fig. 241.—White Mustard, ×110.

Transverse section through mucilaginous epidermis, sub-epidermal parenchyma layer (square cells), palisade cells, and broken parenchyma layer of the hull.

Fig. 242 —White Mustard, XIIO.

Transverse section through the tissue of the radicle.

Fig. 243.—White Mustard X110. Surface view of two layers of the hull or seed coat. Fig. 244 —White Mustard, X110.
Surface section through palisade cells and underlying layer of the seed coat.



## PLATE XXXII.

## SPICES.

Fig. 245.—Black Mustard, ×110.

Transverse section, showing fragments of the epidermis and dark colored palisade cells of the seed coat.

Fig. 246.—Black Mustard, ×110. Surface view of two of the seed coat layers.

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#### PLATE XXXIII.

SPICES.

Fig. 249.—Dakota Mustard Flour, × 110.

Dark spots show starch grains of foreign weed seed, stained with iodine.

Fig. 250.—Adulterated Mustard Flour, × 130. Showing masses of wheat starch.

Fig. 251.—Pepper, ×110.

Transverse section through inner part of pericarp (including parenchyma and seed coat layers) and portion of perisperm, showing starch and oil cells.

Fig. 252.—Pepper, ×110. Surface view of hypodermal layer.



SPICES.

Fig. 253.—Pepper, ×110.

Transverse section through outer part of pericarp, showing epidermis, underlying stone cell layers, parenchyma, and seed coat.

Fig. 254.—Pepper, ×110. Surface section through stone cell layer.

Fig. 255.—Pepper Starch, ×220. Starch granules separated.

Fig. 256.—Pepper Starch, X110. Starch grains in masses.



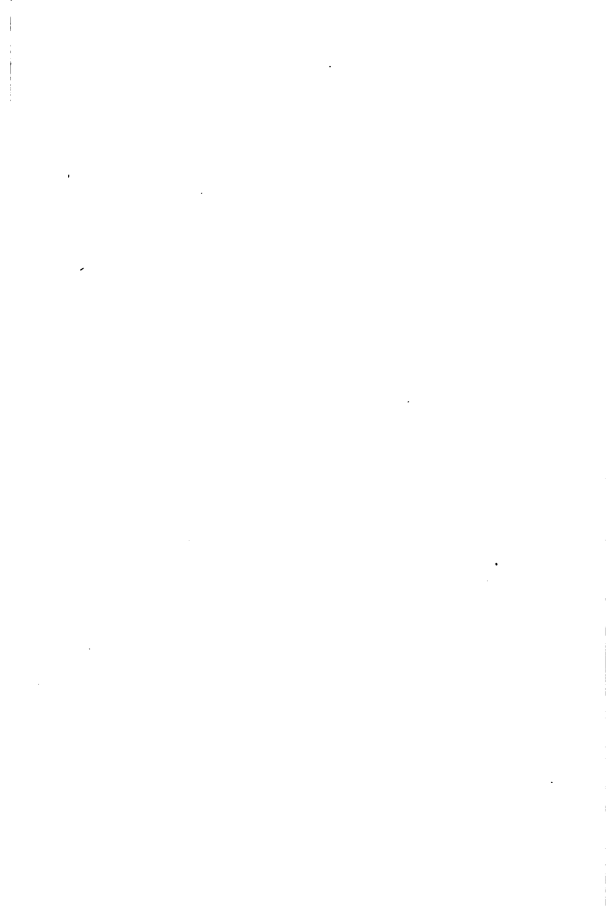
SPICES.

Fig. 257.—Ground Pepper Shells,  $\times$  120. Mainly showing stone cells.

Fig. 258.—Adulterated Pepper, ×130. Showing wheat and buckwheat starches.

FIG 259.—Adulterated Pepper, × 130. Showing wheat, corn, and rice starches.

Fig. 260.—Adulterated Pepper, × 130. The large, lower mass shows buckwheat starch, while the finer-grained mass near the top is of pepper.



## PLATE XXXVI.

# SPICES. SPICE ADULTERANTS.

Fig. 261.—Adulterated Pepper, X110. The central mass shows the sclerenchyma cells of Cayenne and wheat starch are the adulterants. olive stones.

Fig. 262.—Adulterated Pepper,  $\times 130$ .



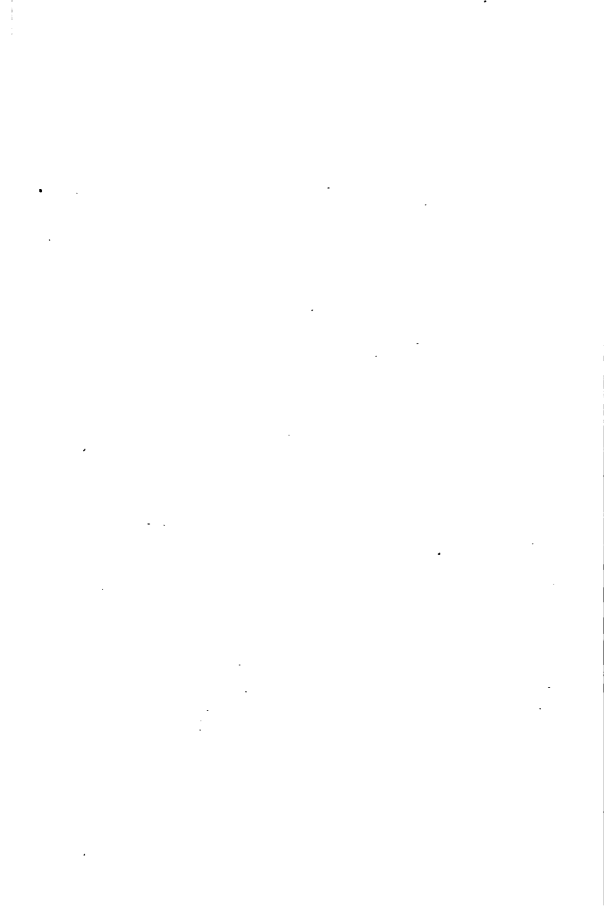
# PLATE XXXVII.

## SPICE ADULTERANTS.

Fig. 265.—Powdered Elm Bark, XIIO.

Fig. 266.—Pine Sawdust, ×110. Finely ground.

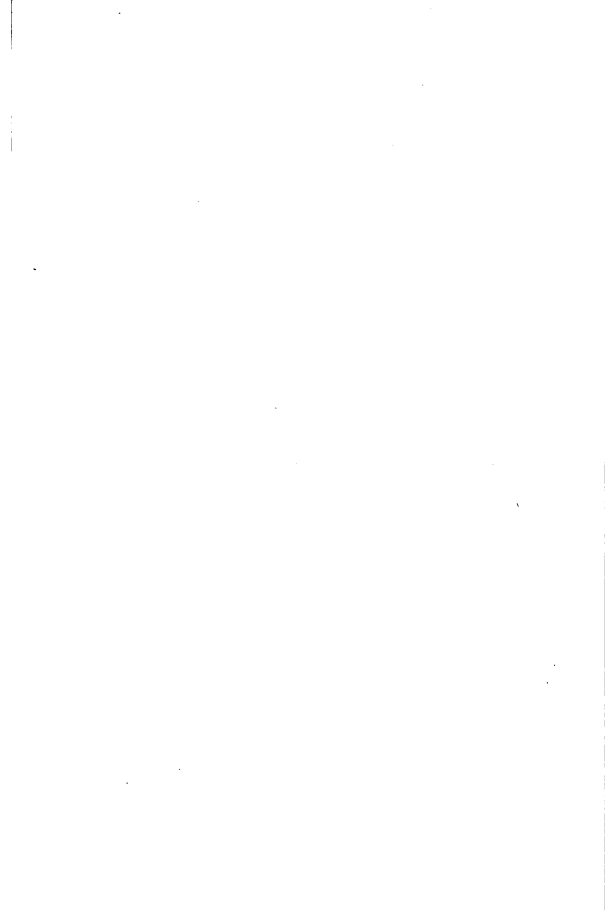
Fig. 267 —Pine Wood, XIIO. Transverse section. Fig. 268.—Pine Wood, ×110. Radial and tangential sections.



## PLATE XXXVIII.

#### EDIBLE FATS.

Ftg. 269.—Pure Butter, ×25.
With polarized light and selenite plate.



## PLATE XXXIX.

#### EDIBLE FATS.

Fig. 272.—Lard Stearin, ×110. Leaf lard, crystallized from ether. Fig. 273.—Lard Stearin, ×220. Leaf lard, crystallized from ether.

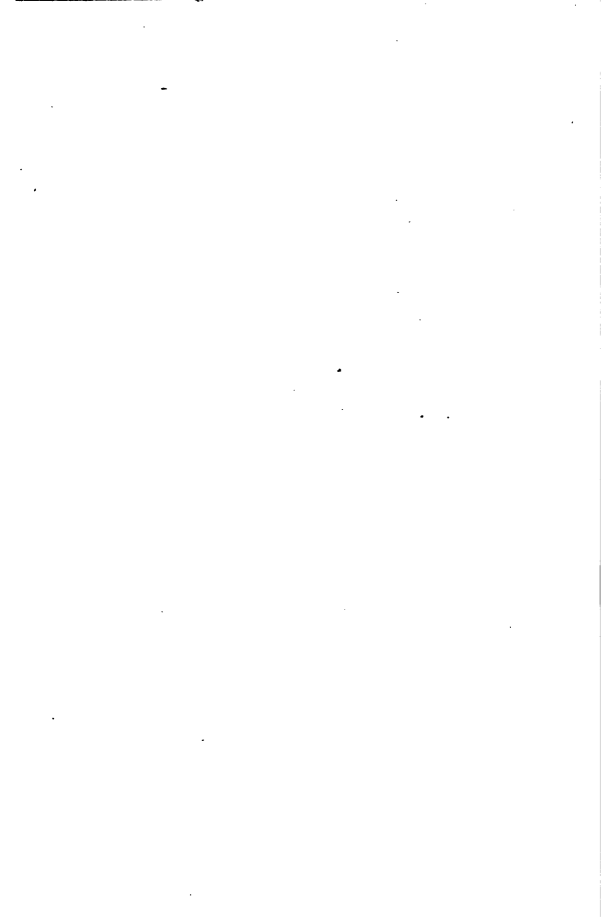


PLATE XL.

EDIBLE FATS.

Fig. 276.—Beef Stearin, ×35. Crystallized from ether.

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